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Medicines used for Multiple Sclerosis – A Health Technology Assessment



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Norwegian Institute of Public Health Oslo, February 2016

Key messages

This Health Technology Assessment was commissioned by the "National system for the introduction of new health technologies within the specialist health service". The aim of this report was to assess the effect and cost-effectiveness of the disease modifying medicines used in Norway for patients with relapsing remitting multiple sclerosis (dimethyl fumarate, terifluno-mide, interferon beta, peg-interferon, glatiramer acetate, natalizumab, fingolimod, and alemtuzumab).

The key results are:

- We identified 37 randomised clinical trials. The quality of the available evidence ranged from very low to high.
- Alemtuzumab 12 mg had the best effect on annual relapse (for medicines we had evidence of high quality). Dimethyl fumarate 240 mg twice daily and fingolimod oral 0.5 mg were the most effective against disability progression (for medicines we had evidence of high quality).
- Our results indicated that interferon beta-1a 44 mcg and peg-interferon beta-1a were associated with more withdrawal due to adverse events than placebo. The examined treatments had no effect on mortality compared to placebo.
- Our health economic analysis, examining all multiple sclerosis treatment alternatives, indicated that alemtuzumab was more effective (in terms of quality-adjusted life-years (QALY)) and less costly than the other treatment alternatives. We did several scenario analyses and the cost-effectiveness results were robust to variations in the model assumptions.
- The results of a scenario analysis that excluded alemtuzumab (the dominant strategy), showed that three treatments alternatives (interferon beta-1b (Extavia), peg-interferon beta-1a and natalizumab) could be cost-effective depending on the willingness-to-pay (WTP) per QALY. Assuming a WTP below NOK 1,000,000, interferon beta-1b

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Health technology assessment (HTA) is a multidisciplinary process that summarizes information about the medical, social, economic and ethical issues related to the use of a health technology in a systematic, transparent, unbiased, robust manner. Its aim is to inform the development of safe, effective health policies that are patient focused and that seek to achieve best value.

Doesn't answer everything:

 Excludes studies that fall outside of the inclusion criteria
 No recommendations

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(Extavia) was 40% likely to be the most cost-effective treatment, followed by peginterferon beta-1a (30% likely).

- The results of our model analysis showed that there is some degree of uncertainty regarding the input parameters. More research on efficacy and epidemiological data would have the greatest impact on reducing decision uncertainty.
- Our bugdet impact analysis based on the results of our cost-effectiveness analysis, the drugs' adverse events profile, and current clinical practice showed that there is a substantial potential for cost saving.

Executive summary

Background

Several disease-modifying therapies are available for the treatment of multiple sclerosis, but the comparative clinical effectiveness of these medicines is unclear. Furthermore, the cost-effectiveness of the different treatments has not been investigated in a Norwegian setting. To ensure the most appropriate multiple sclerosis management, it is important to assess effectiveness and cost-effectiveness of disease modifying medicines used for multiple sclerosis.

Objective

The aim of this project was to compare the effect and cost-effectiveness of the disease modifying medicines used for multiple sclerosis in Norway.

Methods

We conducted a systematic review based on the following conditions: Evidence should come from randomised controlled trials (RCTs) with study populations that included men and women aged 18 years or older were eligible. Modifying medicines used for multiple sclerosis were our intervention of interest (dimethyl fumarate, teriflunomide, interferon beta, peg-interferon, glatiramer acetate, natalizumab, fingolimod, and alemtuzumab). We included studies that compared these medicines to placebo or to each other. We examined the following endpoints: annual relapse, disability progression, mortality, serious adverse events, withdrawal from the study due to adverse events, hospitalisations, and health related quality of life.

We systematically searched the literature for previously published health technology assessment reports or systematic reviews that answered our objectives, and met our inclusion criteria. We conducted a systematic review of randomised controlled trials to supplement the evidence of previously published health technology assessments.

Two persons independently examined the risk of bias of included studies using the Norwegian Knowledge Centre for the Health Services methods. These are based on Cochrane methodology.

4 Executive summary

We summarised the evidence from the randomised clinical trials quantitavely through network meta-analyses of data on direct and indirect evidence on all relevant comparisons.

Two persons independently assessed the quality of the evidence for each selected endpoint. We used GRADE (Grading of recommendations Assessment, Development, and Evaluation) to assess our confidence in the effect estimates.

In order to assess the cost-effectiveness of disease-modifying therapies in patients diagnosed with relapsing-remitting multiple sclerosis, we developed a decision analytic model. The economic model was developed in the form of a cost-utility analysis and included treatments approved and available in Norway. The model structure and all assumptions were adapted to the Norwegian setting based on Norwegian clinical practice. Efficacy estimates were taken from our network meta-analyses. Transitional probabilities were derived from published sources and clinical experts' opinions. Quality of life data were extracted from published studies based on a systematic review of the literature. The costs of medications were based on prices obtained through the Drug procurement cooperation (LIS), and other costs were based on official Norwegian unit prices.

We performed probabilistic sensitivity analyses, designed as a Monte Carlo simulation with 10,000 iterations, to explore the uncertainty surrounding our results.

Results

All examined treatments were more effective than placebo against annual relapse. The effect was best for alemtuzumab 12 mg (based on high quality evidence). Fingolimod oral 0.5 mg and dimethyl fumarate 240 mg twice daily were also associated with a reduction in annualised relapse rate. For disability progression, dimethyl fumarate 240 mg twice daily and fingolimod 0.5 mg were more effective than placebo (high quality evidence).

For withdrawal due to adverse events, the conclusion is unclear due to the low quality of the available evidence. However, our results indicate that interferon beta-1a 44 mcg, and peg-interferon beta-1a are associated with more withdrawal due to adverse events than placebo.

For the outcomes change in expanded disability status scale, serious adverse events, and mortality; we did not assess the quality of the available evidence. Our results indicate that interferon beta-1a 30 mcg is associated with a reduction in expanded disability status scale. Interferon beta-1a 30 mcg is associated with fewer serious adverse

events. Finally, our results showed that none of the examined treatments increased or decreased mortality compared to placebo.

Our health economic analysis indicated that alemtuzumab dominated all other disease-modifying therapies, as it was more effective in terms of quality-adjusted lifeyears (QALY) and less costly than the other treatment alternatives.

A scenario analysis that excluded alemtuzumab (the dominant strategy) showed that three treatment alternatives (interferon beta-1b (Extavia), peg-interferon beta-1a and natalizumab) could be cost-effective depending on the willingness-to-pay (WTP) threshold. Interferon beta-1b was likely to be the cost-effective choice for a WTP per QALY below NOK 1,658,000. Peg-interferon was the cost-effective option for a WTP from NOK 1,658,450 to NOK 10,619,960, and natalizumab was the cost-effective alternative for a WTP above NOK 10,619,960. Assuming a WTP below NOK 1,000,000 per QALY, interferon beta-1b (Extavia) was approximately 40% likely to be the most cost-effective treatment, followed by peg-interferon beta-1a (approximately 30% likely).

The results of probabilistic analysis showed that there is some degree of uncertainty regarding the input parameters. More research on efficacy and epidemiologic input parameters would have the greatest impact on reducing decision uncertainty.

We performed several scenario analyses to test the uncertainty around the model assumptions. The results showed that, while there were numerical changes to the incremental cost-effectiveness ratio, the cost-effectiveness results were robust to variations in the model assumptions and the conclusions of the analysis would not change.

Our bugdet impact analysis based on the results of our cost-effectiveness analysis, the drugs' adverse events profile, and current clinical practice showed that there is a substantial potential for cost saving.

Discussion

We used a systematic methodology to search for evidence, extract data, and assess the risk of bias of studies and quality of evidence for important outcomes. The systematic review included evidence on both established and emerging treatments. We examined the effect of these treatments on clinical endpoints relevant for patients with multiple sclerosis. We have analysed direct and indirect evidence through network meta-analyses. The consistency of results using different methods indicates that our results are robust.

Our systematic review has some limitations, due more to the weakness of the available evidence than to the methods used in this report. These limitations are related to the

paucity and quality of the available evidence, and to the methodologies used in the included randomised controlled trials.

We used a probabilistic Markov-model, considered the appropriate approach for simulating the natural history of multiple sclerosis. The model structure and all assumptions have been adapted to the Norwegian setting based on Norwegian clinical practice with close assistance of experts in this field.

For transitional probabilities, we did not find Norwegian data that were compatible with the developed model, so these were based on estimates reported in the published literature.

Study designs of published trials did not permit separate analyses of first and second line treatments, or conclusions regarding the sequential use of first and second line treatments. Therefore, we did not perform separate cost-effectiveness analyses for first or second line treatments. In addition, based on expert opinion, we did not include combination therapy in our model, as it is not relevant to current Norwegian clinical practice.

Conclusion

Alemtuzumab 12 mg had the best effect against annual relapse. Dimethyl fumarate 240 mg twice daily and fingolimod oral 0.5 mg were the most effective against disability progression. Results indicate that some treatments are associated with more withdrawals due to adverse events than placebo. Our results showed that the examined treatments had no effect on mortality.

Our health economic analysis indicated that alemtuzumab was more effective and less costly than the other treatment alternatives. A scenario analysis that excluded alemtuzumab indicated that three treatment alternatives (interferon beta-1b (Extavia), peg-interferon beta-1a and natalizumab) could be cost-effective depending on the WTP. For a WTP below NOK 1,000,000 per QALY, interferon beta-1b (Extavia) was approximately 40% likely to be the most cost-effective treatment, followed by peg-interferon beta-1a (approximately 30% likely).

The results of probabilistic analysis showed that there is some degree of uncertainty regarding the input parameters. More research on efficacy and epidemiologic input parameters would have the greatest impact on reducing decision uncertainty.

Our budget impact analysis showed that there is a substantial potential for cost saving.

Hovedfunn (norsk)

Denne fullstendige metodevurderingen ble bestilt av «Nasjonalt system for innføring av nye metoder i spesialisthelsetjenesten». Målet var å sammenligne effekt, sikkerhet og kostnadseffektivitet av sykdomsmodifiserende legemidler som brukes for multippel sklerose i Norge (dimetylfumarat, teriflunomid, interferon beta, peginterferon, glatirameracetat, natalizumab, fingolimod og alemtuzumab).

Hovedfunnene er:

- Vi identifiserte 37 randomiserte kontrollerte studier og kvaliteten på dokumentasjon varierte fra veldig lav til høy.
- Basert på sammenligninger hvor kvaliteten på dokumentasjonen var høy kan vi si at alemtuzumab 12 mg hadde den beste effekten mot årlig tilbakefall, og at dimetylfumarat 240 mg to ganger om dagen og fingolimod 0.5 mg var de mest effektive mot sykdomsprogresjon.
- Våre resultater indikerer at interferon beta-1a 44 mcg, og peginterferon beta-1a var assosiert med høyere frafall på grunn av bivirkninger enn placebo. Våre resultater viste ingen av behandlingene hadde effekt på dødelighet.
- Vår helseøkonomiske analyse indikerte at alemtuzumab var bedre og mindre kostnadskrevende enn de andre behandlingsalternativene. Vi utførte flere scenarioanalyser for å teste usikkerheten rundt forutsetninger ved modellen, men konklusjonene endret seg ikke.
- En scenarioanalyse hvor alemtuzumab (den dominante strategien) ble ekskludert, viste at tre behandlingsalternativer (interferon beta-1b (Extavia), peginterferon beta-1a og natalizumab) kunne være kostnadseffektive, avhengig av betalingsvilje per vunnet kvalitetjustert leveår (QALY). Ved å anta en betalingsvilje under en million kroner per vunnet QALY, var interferon beta-1b (Extavia) trolig den mest kostnadseffektive behandlingen (ca. 40 %), etterfulgt av peginterferon beta-1a (ca. 30 %).

Tittel:

Fullstendig metodevurdering av legemidler ved multippel sklerose

Publikasjonstype: Metodevurdering

En metodevurdering er resultatet av å

- innhente
- kritisk vurdere og

- sammenfatte relevante forskningsresultater ved hjelp av forhåndsdefinerte og eksplisitte metoder.

Minst ett av følgende tillegg er også med:

helseøkonomisk evaluering, vurdering av konsekvenser for etikk, jus, organisasjon eller sosiale forhold

Svarer ikke på alt:

Ingen studier utenfor de eksplisitte inklusjonskriterieneIngen anbefalinger

Hvem står bak denne rapporten?

Folkhelseinstuttet har skrevet rapporten på oppdrag fra Nasjonalt system for innføring av nye metoder i spesialisthelsetjenesten

Når ble litteratursøket utført?

Søk etter studier ble avsluttet november 2015

- Vår modellanalyse viste at det er en viss grad av usikkerhet knyttet til parameterne brukt i analysen. Mer forskning på effekt av legemidlene eller bedre epidemiologiske data fra norske registre ville hatt størst innvirkning på å redusere beslutningsusikkerhet.
- Vår budsjettkonsensanalyse basert på resultatene av vår kostnadseffektivitetsanalyse, bivirkninger knyttet til behandlingsalternativene og dagens kliniske praksis viste at det er et betydelig potensial for å redusere kostnadene knyttet til MS-behandling i spesialisthelsetjenesten.

Sammendrag (norsk)

Fullstendig metodevurdering av legemidler ved multippel sklerose

Bakgrunn

Det finnes flere sykdomsmodifiserende legemidler godkjent til bruk ved multippel sklerose, men en fullstendig sammenligning av den kliniske effektiviteten på tvers av alle disse har ikke vært gjort. Kostnadseffektiviteten av de ulike behandlingene er heller ikke blitt undersøkt i en norsk setting.

Problemstilling

Målet vårt var å sammenligne effekt, sikkerhet og kostnadseffektivitet av sykdomsmodifiserende legemidler som brukes for multippel sklerose i Norge.

Metode

Vi utførte en systematisk oversikt, hvor vi inkluderte randomiserte kontrollerte studier på personer over 18 år med multippel sklerose behandlet med følgende legemidler: dimetylfumarat, teriflunomid, interferon beta, peginterferon, glatirameracetat, natalizumab, fingolimod og alemtuzumab. Vi inkluderte studier som sammenlignet disse medisinene med placebo eller med hverandre. Vi undersøkte følgende kliniske endepunkt: årlig attakk, sykdomsprogresjon, dødelighet, alvorlige bivirkninger, frafall fra studien på grunn av bivirkninger, sykehusinnleggelser og helserelatert livskvalitet.

Vi søkte etter publiserte Health Technology Assessment (HTA) rapporter og systematiske oversikter som besvarte vår problemstilling. Deretter søkte vi etter randomiserte kontrollerte studier for å supplere kunnskapsgrunnlaget med informasjon publisert etter søkedato i den nyeste, mest omfattende HTA rapporten vi identifiserte.

To personer undersøkte uavhengig av hverandre kvaliteten på den inkluderte HTArapporten og risiko for systematiske skjevheter i de supplerende studiene. Vi oppsummerte kliniske resultater gjennom nettverks meta-analyser som baserer seg på både direkte og indirekte sammenligninger. Til slutt brukte vi GRADE (Grading av anbefalinger Assessment, Development, and Evaluation) for å vurdere kvaliteten på dokumentasjonen og vår vår tillit til effektestimatene.

For å vurdere kostnadseffektiviteten av de sykdomsmodifiserende legemidlene hos pasienter med relapsing-remitting multippel sklerose, utviklet vi en helseøkonomisk modell (Markov-modell). Modellstruktur og alle forutsetninger ble tilpasset norsk klinisk praksis. Effektestimatene ble tatt fra vår systematiske gjennomgang av klinisk effekt og sikkerhet. Overgangssannsynligheter ble hentet fra publiserte kilder og supplert med opplysninger fra kliniske eksperter. Livskvalitetsdata ble hentet fra publiserte studier indentifisert gjennom en systematisk gjennomgang av litteratur. Kostnader på medisiner ble basert på priser fra Legemiddelinnkjøpssamarbeidet (LIS), og andre kostnader var basert på norske kilder. Vi utførte probabilistiske sensitivitetsanalyser, utformet som en Monte Carlo-simulering med 10,000 gjentakelser, for å analysere usikkerheten i våre resultater.

Resultat

Alle undersøkte legemidler var mer effektive enn placebo mot årlig attakk. Effekten var best for alemtuzumab 12 mg (basert på evidens av høy kvalitet). For sykdomsprogresjon var dimetylfumarat og fingolimod mer effektivt enn placebo (evidens av høy kvalitet).

For frafall på grunn av bivirkninger var det lavere kvalitet på tilgjengelig dokumentasjon, noe som knytter mer usikkerhet til resultatene. Men våre resultater indikerer at både interferon beta-1a 44 mcg, og peginterferon beta-1a begge er assosiert med høyere frafall på grunn av bivirkninger enn placebo.

Vi vurderte ikke kvaliteten på tilgjengelig dokumentasjon om endring i uførhetsstatusskalaen EDSS (expanded disabliltity symptom scale), alvorlige bivirkninger og dødsfall. Våre resultater tyder på at interferon beta-1a 30 mcg var relatert til en reduksjon i EDSS nivå. Interferon beta-1a 30 mcg var assosiert med færre alvorlige bivirkninger. Til slutt, viser våre resultater at ingen av de undersøkte behandlinger ga økt dødelighet sammenlignet med placebo.

Vår helseøkonomiske analyse indikerte at alemtuzumab dominerte alle andre sykdomsmodifiserende behandlinger. Alemtuzumab var både mer effektiv og mindre kostnadskrevende enn de andre behandlingsalternativene.

Resultatene av en scenarioanalyse hvor alemtuzumab (den dominante strategien) ble ekskludert, viste at tre behandlingsalternativer (interferon beta-1b (Extavia), peginterferon beta-1a og natalizumab) kunne være kostnadseffektive, avhengig av betalingsvilje per vunnet kvalitetjustert leveår (quality-adjusted life-years, QALY). Forutsatt en betalingsvilje (Willingness to pay, WTP) lavere enn 1658 000 kroner per QALY, vil Interferon beta-1b sannsynligvis være et kostnadseffektivt valg. For en WTP mellom 1 658 450 og 10 619 960 kroner var peginterferon et kostnadseffektivt alternativ, og for en WTP over 10 619 960 kroner var natalizumab et kostnadseffektivt alternativ. Ved å anta en betalingsvilje på under 1 000 000 kroner per vunnet QALY var interferon beta-1b (Extavia) trolig den mest kostnadseffektive behandlingen (ca. 40%), fulgt av peginterferon beta-1a (ca. 30%).

Sannsynlighetsanalyser viste at det er usikkerhet knyttet til parameterne benyttet i modellen. Mer forskning på effekt av legemidlene eller bedre epidemiologiske data fra norske registre ville hatt størst innvirkning på å redusere beslutningsusikkerhet.

Vi utførte flere scenarioanalyser for å teste usikkerheten rundt ulike helseøkonmiske modellforutsetninger. Selv om det var numeriske endringer i resultater, så var resultatene for kostnadseffektivitet robuste og konklusjonene fra analysen endret seg ikke.

Vår budsjettkonsensanalyse basert på resultater av kostnadseffektivitetsanalysen vår, bivirkninger knyttet til behandlingsalternativene og dagens kliniske praksis viste at det er et betydelig potensial for å redusere kostnadene knyttet til MS-behandling i spesialisthelsetjenesten.

Diskusjon

Vi brukte internasjonalt anerkjente metoder for å systematisk oppsummere kunnskapsgrunnlaget og fokuserte på kliniske endepunkter som er relevante for pasienter med multippel sklerose. Konsistente resultater ved bruk av direkte, indirekte eller nettverksanalyser viser at våre resultater er pålitelige.

Vår systematiske gjennomgang har noen begrensninger. De er hovedsakelig knyttet til at det er få studier eller rapporterte utfall for enkelte av sammenligningene og metodiske uklarheter i de inkluderte randomiserte kontrollerte studiene.

Vi brukte en probabilistisk Markov-modell, som er ansett for å være den beste måten å simulere sykdomsforløpet til multippel sklerose på. Modellens struktur og alle forutsetninger er tilpasset norske forhold og klinisk praksis med tett bistand fra eksperter på feltet. Der vi ikke fant norske data som kunne brukes i modellen benyttet vi overgangssannsynligheter fra publisert litteratur.

Måten de publiserte kliniske studiene er utført på gjør det vanskelig å undersøke første- og andrelinje behandlinger hver for seg, eller å konkludere på sekvensiell bruk av ulike behandlinger. Vi utførte derfor ikke separate kostnadseffektivitetsanalyser for første- eller andrelinjebehandlinger. Som følge av ekspertuttalelser, gjorde vi heller ikke analyser for kombinasjonsbehandling siden det ikke er relevant for norsk klinisk praksis i dag.

Konklusjon

Basert på dokumentasjon av høy kvalitet kan vi si at alemtuzumab 12 mg hadde den beste effekten mot årlig tilbakefall og at fingolimod oral 0,5 mg og dimetylfumarat 240 mg to ganger daglig hadde den beste effekten mot sykdomsprogresjon. Resultatene tyder på at noen behandlinger er forbundet med mer frafall på grunn av bivirkninger enn placebo. De inkluderte intervensjonene hadde ingen effekt på dødelighet.

Vår helseøkonomiske analyse indikerte at alemtuzumab var både mer effektiv og mindre kostnadskrevende enn de andre behandlingsalternativene.

En scenarioanalyse hvor alemtuzumab ble ekskludert viste at tre behandlingsalternativer (interferon beta-1b (Extavia), peginterferon beta-1a og natalizumab) kunne være kostnadseffektive, avhengig av betalingsvilje per vunnet QALY. Ved å anta en betalingsvilje under en million kroner per vunnet QALY, var trolig interferon beta-1b (Extavia) den mest kostnadseffektive behandlingen (ca. 40 %), fulgt av peginterferon beta-1a (ca. 30 %).

Resultatene av sannsynlighetsanalysen viste at det er en viss grad av usikkerhet knyttet til de ulike parameterne inkludert i analysen. Mer forskning på effekt og epidemiologiske data vil ha størst innvirkning på å redusere usikkerheten rundt beslutningen.

Vår budsjettkonsensanalyse viste at det er et betydelig potensial for å redusere kostnadene knyttet til MS-behandling i spesialisthelsetjenesten.

Glossary and abbreviations	
CI	Confidence interval. A measure of uncertainty around the results of a statistical analysis that describes the range of values within which we can be reasonably sure that the true mean effect lies. Wider intervals indicate lower precision; narrow intervals, greater precision.
CIS	Clinical isolated syndrome
CNS	Central nervous system
CUA	Cost-utility analysis. An economic evaluation where health consequences are measured in QALY s.
EDSS	Expanded disability status scale
EQ-5D	European Quality of Life-5 Dimensions. EQ-5D is a standardized instrument for use as a measure of health outcome.
EVPI	Expected value of partial perfect information
GRADE	Grading of recommendations Assessment, Development, and Evaluation
НТА	Health Technology Assessment
Healthcare perspective	Economic evaluation from a healthcare perspective will consider only the costs and consequences specifically related to the healthcare sector (direct costs), <i>e.g.</i> staff costs, capital costs, drug acquisition costs.
ICER	Incremental cost-effectiveness ratio . The ratio of the difference in costs between two alternative health technologies to the difference in effectiveness between these two technologies. $ICER = \frac{Cost_{\text{intervention}} - Cost_{\text{comparator}}}{Effect_{\text{intervention}} - Effect_{\text{comparator}}} = \frac{\Delta C}{\Delta E}$
MRI	Magnetic resonance imaging
MS	Multiple sclerosis
NHB	Net Health Benefit. In a decision-making process, a positive NHB suggests that the intervention represents good value for money $NHB = \Delta E - \frac{\Delta C}{\lambda}$
NMB	Net Monetary Benefit. In a decision-making process, a positive NMB suggests that the intervention represents good value for money. $NMB = \lambda \cdot \Delta E - \Delta C$
Odds	The odds of an event happening is defined as the probability that an event will occur divided by the probability that the event will not occur.

OR	Odds ratio. The ratio of the odds of an outcome in one treatment group divided by the odds of the same outcome in a different treatment group.
PPMS	Primary progressive multiple sclerosis
PSA	Probabilistic sensitivity analysis. An analysis of the uncertainty re- lated to all parameters in a decision analytic model. Typically performed by Monte Carlo simulation, hence by drawing values from probability distributions for all parameters simultaneously
QALY	Quality-adjusted life-year. A measure of health outcomes that combines quantity and quality of life by assigning to each year of life a weight from 1 (perfect health) to 0 (state judged equivalent to death) dependent on the individual's health related quality of life during that year
RCT	Randomised controlled trial. An experiment in which investigators use randomisation to allocate participants into the groups that are being compared. Usually allocation is made at the level of individuals, but sometimes it is done at group level e.g. by schools or clinics. This design allows assessment of the relative effects of interventions.
RRMS	Relapsing-remitting multiple sclerosis
RR	Relative risk / risk ratio. The relative risk is the absolute risk (AR) in the intervention group divided by the AR in the control group. It is to be distinguished from odds ratio (OR), which is the ratio of events over non-events in the intervention group over the ratio of events over non-events in the control group.
SPMS	Secondary progressive multiple sclerosis
SR	Systematic review. A review of a clearly formulated question that uses systematic and explicit methods to identify, select, and critically appraise relevant research, and to collect and analyse data from the studies that are included in the review. Statistical methods (meta-analysis) may or may not be used to analyse and summarise the results of the included studies.
Statistically significant	Means that the findings of a study are unlikely to have arisen because of chance. Significance at the commonly cited 5% level ($P < 0.05$) means that the observed difference or greater difference would occur by chance in only 1/20 similar cases. Where the word "significant" or "significance" is used without qualification in the text, it is being used in this statistical sense.
SUCRA	Surface under the cumulative ranking curve
WTP (λ)	Willingness to pay . A pre-specified limit of what society is willing to pay for a given health unit (e.g. QALY or life year). In Norway it is common to use NOK 500 000 per QALY or life year in economic evaluations.

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Preface

This project was commissioned by the "National system for the introduction of new health technologies within the specialist health service", that wanted us to examine the effect and cost-utility of the disease modifying medicines used for patients with relapsing remitting multiple sclerosis in Norway. The results will be used as scientific documentation for price negotiations, and guidelines development.

Elisabeth Couto was lead reviewer for the clinical evaluation and Vida Hamidi led the health economic evaluation. Rune Midgard and Torbjørn Wisløff performed peer review of the report. We thank Elisabeth Gulowsen Celius and Elisabeth Farbu for clinical expertise and input in the report, and Bjørn Svendsen for his contribution on cost information.

The project group consisted of:

- Couto, Elisabeth, (project leader), senior researcher
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We would like to thank Elisabeth Gulowsen Celius, Elisabeth Farbu, Rune Midgard, Bjørn Svendsen, and Torbjørn Wisløff for their expertise in this project. The Norwegian Institute of Public Health assumes final responsibility for the content of this report.

The aim of this report is to support well-informed decisions in health care that lead to improved quality of services. The evidence should be considered together with other relevant issues, such as clinical experience and patients' preferences.

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Objective

Overall objective

• To examine the effect and cost-utility of the disease modifying medicines used for patients with relapsing remitting multiple sclerosis in Norway.

Specific objectives

- To conduct a systematic review to assess the efficacy and safety of the different disease modifying medicines used for multiple sclerosis with regard to clinical important endpoints
- To carry out a health economic evaluation ascertaining cost-utility of the disease modifying medicines used for patients with relapsing remitting multiple sclerosis.

Background

Multiple sclerosis (MS) is a chronic inflammatory disorder of the central nervous system (CNS) with secondary neurodegeneration (1). It affects nerves in the brain and spinal cord by damaging the myelin sheath that covers the axon part of the nerve cells. The myelin sheath protects and aids signal transduction, therefore, when damaged, it affects the transfer of timely and correct information from the CNS to the peripheral part of the nervous system (1-3).

The epidemiology of multiple sclerosis

MS is one of the most common causes of disability in young adults (4). In 2013, a systematic review summarised MS incidence and prevalence estimates reported by 123 studies that used a range of different data sources (5). Prevalence and incidence estimates tended to be higher in Northern countries, and in more recently published studies. Incidence surveys show an increase in MS incidence in later years (6). Reported annual incidence rates are 1.9 (95% confidence interval: 1.2-2.6) for the period 1953 to 1957, and 8.5 (7.3.9.7) for 1978 to 2007 (7). Increase in MS incidence could be due, to some extent, to changes in methods and criteria used for MS diagnosis (6). In Europe, the prevalence of MS is twice as high in women than in men (5). Incidence rates are generally also higher in women (5). A study, using data from the National Patient Registry, the Norwegian MS registry, and Biobank data estimated crude prevalence rates of 203/100,000 (95% confidence interval 199 – 207) overall, 280 (247-287) for women, and 126 (122-130) for men (8).

The disease usually starts around the age of 30 (range 20-40), and prevalence rates peak at around 50 (6). The median time to death is around 30 years from disease onset, representing a reduction in life expectancy of 5 to 10 years (1).

The aetiology of MS is not well understood. Geographical variations in MS prevalence and incidence could be due to differences in genes and environment. To date, most commonly reported risk factors for MS are exposure to Epstein Barr virus, cigarette smoking, low sunlight exposure and vitamin D levels and genetic predisposition (1, 9-11).

The clinical course and diagnosis of multiple sclerosis

Clinical manifestations depend on the affected area of the CNS. Symptoms reflect an involvement of motor, sensory, visual and autonomic systems (1). Symptoms evolve over time. MS appears in several degrees of severity from a mild form (with few attacks) to a more progressive disease that is potentially highly disabling and that impacts on the quality of life of patients and their families (1, 12).

Appropriate MS diagnosis allows early disease management. Different diagnosis criteria have been used over the years, leading to possible differences in MS diagnosis with time. The revised McDonald criteria are the most commonly used for MS diagnosis nowadays. National guidelines, such as British (NICE) and Norwegian guidelines, recommend the use of the revised McDonald criteria for MS diagnosis (12, 13).

To be diagnosed with MS, patients should have at least one clinical attack (demyelinating event in the CNS with duration of symptoms of more than 24 hours in the absence of fever or infection) corroborated by findings on neurological examination, visual evoked potential response or findings on magnetic resonance imaging (MRI) consistent with demyelination in the CNS (T2 lesion or T1 gadolinium-enhancing lesion). In addition, exclusion of other possible diagnoses is essential for the diagnosis of MS.

MS is classified as (1, 13):

- Clinical isolated syndrome (CIS): one attack and objective clinical evidence of one lesion.
- Relapsing-remitting MS (RRMS): objectively established disease as with two or more clinical attacks and localisaton of two or more lesions in the CNS. It is characterised by episodes of acute worsening of function followed by partial or complete recovery (14). 85 to 90% of patients present with RRMS (11). Aproximately half of the patients with RRMS will develop secondary progressive MS (15).
- Secondary progressive MS (SPMS): About 30-40% of the prevalent MS population have SPMS. It is associated with disease progression without clinical attacks and of highly variable degrees (16).
- Primary progressive MS (PPMS): at least one year of disease progression and characteristic findings on MRI and/or positive findings in cerebrospinal fluid.

Disease progression is most commonly assessed by relapse rate and disease progression. The gradual increasing level of disability is often measured with the Expanded Disability Status Scale (EDSS), an ordinal scale ranging from 0 (normal clinical status) to 10 (death due to MS) in steps of 0.5 points (17).

Treatment alternatives

Disease-modifying medicines are the standard treatment for patients with MS. It is possible to treat both the underlying disease, relapses and MS-related symptoms. Disease modifying drugs may inhibit the inflammatory process to prevent progression and reduce disabilities due to the disease. The different treatment options have different mechanisms of action, routes of administration, approved indications and other differences influencing their use. The various medications are presented in Table 1.

Due to safety issues, some of these treatments are used as first line treatments (dimethyl fumarate, teriflunomide, interferon beta, peg-interferon, glatiramer acetate), and others as second line treatments (natalizumab, fingolimod, and alemtuzumab) according to different national guidelines (18).

Disease-modifying treatments are expensive. The use of MS medicines has been described as "uneven" with "questionable effects on the long-term accumulation of disability and disease progression" (1). Currently a number of new disease-modifying therapies are available for the treatment of MS, but it is uncertain whether the new medicines are cost-effective in the Norwegian setting. To insure proper MS management, it is important to assess the effectiveness and cost-effectiveness of disease modifying medicines used for MS.

This report was ordered by the "National system for the introduction of new health technologies within the specialist health service", and will be used for price negotiations and guidelines development.

Intervention	Administration form and	Approved indication
Medication name	recommended dose	
First authorisation date in Norway		
Alemtuzumab	- 12 mg concentrate for solution for infusion	Adult patients with relapsing remitting multiple sclerosis
(Lemtrada)	- 12 mg/day for 5 consecutive days, then after 12	(RRMS) with active disease defined by clinical or imaging
Sept.2013	months: 12 mg/day for 3 consecutive days.	features
	Diluted and i.v. over approximately 4 hours	
Dimethyl fumarate	-120 or 240 mg gastro-resistant hard capsules	Adult patients with relapsing remitting multiple sclerosis
(Tecfidera)	- 240 mg twice daily	
Jan. 2014		
Fingolimod	- 0,5 mg hard capsules	- High disease activity despite treatment with at least one
(Gilenya)		disease modifying therapy
March 2011	- 0,5 mg once daily	- Rapidly evolving severe relapsing remitting multiple
		sclerosis
Glatiramer acetat	- 20 mg/ml Solution for Injection, Pre-filled	- Patients experienced a well-defined first clinical episode,
(Copaxone)	Syringe	determined to be at high risk of developing clinically definite
Februar 2004	- 20 mg of glatiramer acetate (one pre-filled	multiple sclerosis
April 2015 (40 mg)	syringe), administered as a subcutaneous	- Ambulatory patients with relapsing, remitting multiple
	injection once daily	sclerosis w/ \geq 2 attacks of neurological dysfunction over the
	- 40 mg of glatiramer acetate administered	preceding two-year period.
	three times weekly	
Interferon beta-1a	- 30 micrograms (6 million IU) powder and	-Relapsing multiple sclerosis w/ ≥ 2 relapses in the previous
(Avonex)	solvent for solution for injection	three years without evidence of continuous progression
May 2011	- 30 micrograms (1 ml solution), by	between relapses
	intramuscular (IM) injection once a week	

Table 1. Overview of included interventions

Interferon beta-1a	- 22 micrograms (6 million IU) solution for	Relapsing multiple sclerosis, w/ ≥ 2 acute exacerbations in the
(Rebif)	injection in pre-filled syringe	previous two years
June 2010	- 44 micrograms given three times per week by	
	subcutaneous injection	
Peg-interferon beta-1a	- 125 micrograms injected subcutaneously every	Adult patients for the treatment of relapsing remitting
(Plegridy)	2 weeks	multiple sclerosis
July 2014		
Interferon beta-1b	- 250 microgram (8.0 million IU) /ml, powder	- Patients with a single demyelinating event with an active
(Betaferon)	and solvent for solution for injection [300	inflammatory process ()determined to be at high risk of
August 2008	microgram (9.6 million IU) per vial]	developing clinically definite multiple sclerosis
	- 250 microgram (8.0 million IU), contained in	- Patients with relapsing-remitting multiple sclerosis w/ ≥ 2
	1 ml of the reconstituted solution, to be injected	relapses within the last two years
	subcutaneously every other day	-Patients with secondary progressive multiple sclerosis with
		active disease, evidenced by relapses.
Interferon beta-1b	See: interferon beta-1b (Betaferon) above	Adults and adolescents from 12-17 years of age.
(Extavia)		Indication similar to Interferon beta-1b (Betaferon) above
June 2006		
Natalizumab	- 300 mg concentrate for solution for infusion	- Adult patients with relapsing remitting multiple sclerosis
(Tysabri)	- 300 mg by i.v over approximately 1 hour, once	-High disease activity despite treatment with a
June 2006	every 4 weeks	betainterferon or glatiramer acetate
		-Rapidly evolving severe relapsing remitting multiple
		sclerosis
Teriflunomide	- 14 mg film-coated tablets	Adult patients with relapsing remitting multiple sclerosis
(Aubagio)	- 14 mg once daily, swallowed whole with some	
Aug.2013	water	

Introduction to Economic Evaluations of Health Care Programmes

The basic task of any economic evaluation is to identify, measure and compare costs and consequences of the alternatives under consideration in an incremental analysis—one in which the differences in costs are compared with differences in consequences (19). Results of economic evaluations can be expressed as an incremental cost-effectiveness ratio (ICER), which is defined by the following equation:

$$ICER = \frac{Cost_{intervention} - Cost_{comparator}}{Effect_{intervention} - Effect_{comparator}} = \frac{\Delta C}{\Delta E}$$

Because the health care sector, like the society in general, is restricted by scarce resources and budget constraints, economic evaluations are important tools for decision makers facing questions of how to prioritize treatments and maximize health benefits using scarce resources. For an economic evaluation to be meaningful in a decision making process, the ICER must be judged with regard to a ceiling ratio that reflects the decision maker's maximum willingness to pay (WTP) for a health gain. The decision rule for an economic evaluation can therefore be expressed as

$$\frac{\Delta C}{\Delta E} < \lambda$$

where λ equals WTP, and means that if the ICER of an intervention is below the ceiling ratio, introducing the intervention represents good value for money. Because the ICER has poor statistical properties, ICERs are often rearranged to express either incremental net monetary benefit (INMB) or incremental net health benefit (INHB), which yields the following decision rules related to INMB or INHB.

INMB: $\lambda \cdot \Delta E - \Delta C > o$ INHB: $\Delta E - (\Delta C/\lambda) > o$

An intervention can in other words be considered cost-effective if it yields a positive INHB or INMB.

Economic evaluations are often based on decision models (such as decision trees, Markov models, etc.) that calculate results based on various input parameters in the model. Because there are always uncertainties related to the values of these parameters, sensitivity analysis is an important feature of any economic evaluation based on a decision model framework. In short, sensitivity analysis illustrates how much the results vary when model parameters are changed. Probabilistic sensitivity analysis (PSA) is a kind of sensitivity analysis. The advantage of PSA is that it makes it possible to take the uncertainties of all of the model-parameters into account simultaneously. The basic approach in PSA is to assign appropriate probability distributions to the model-parameters, which makes it possible to replace the "fixed" values of the parameters with values generated by random draws from the distributions. Doing this repeatedly, with a specified number of iterations, makes it possible to estimate the probabilities that alternative interventions are cost-effective subject to different ceiling values of WTP. The calculation is based on the alternative that renders the highest values of NMB or NHB. Results from PSAs are often presented as scatter plots, which show point estimates of the ICER for all iterations in the cost-effectiveness plane, and also as cost-effectiveness acceptability curves (CEACs), which show the probability of the alternatives being cost-effective subject to changing values of WTP.

Another result from PSA is the expected value of perfect information (EVPI). This number indicates the value to society to have more accurate information about the decision, given a WTP. If EVPI for a given population seems large, it might be of interest to determine for which parameters it would be most useful to obtain additional data. Expected value of perfect information for parameters is a more time-consuming analysis that can help determine for which single parameters or groups of parameters it is most cost-effective to conduct new research.

In short, making a model probabilistic means that it is possible to estimate the uncertainty associated with a decision to implement alternative interventions, and it provides a possibility of estimating the value of collecting additional information from new research.

Priority setting criteria

According to Norwegian policy documents (20) , a treatment should be prioritized if the following criteria are met:

- *The disease is severe:* A disease is considered severe to the degree that it causes pain and discomfort, loss of physical, psychological and social function and if it limits the individual in his or her daily activities. Severity is also evaluated according to the risk increase the disease entails in terms of death, disability and discomfort, if treatment is postponed.
- *The treatment is effective:* The patient should be expected to benefit from treatment in terms of longevity or improved quality of life of certain duration. The treatment effectiveness should also be well documented.
- *The treatment is cost-effective:* The additional costs of the treatment should be reasonable compared to the additional benefits.

It should be mentioned that there is no academic or political consensus regarding what constitutes a reasonable relationship between incremental costs and effects in Norway. For this reason, we use a range of potential willingness-to-pay (WTP) values throughout our report.

Clinical evaluation – Methods

Criteria for considering studies for this review

Type of studies

We searched for published health technology assessment (HTA) reports or systematic reviews (SR) of randomised controlled trials (RCT). We included only reports and reviews of high quality that fitted our inclusion criteria. We supplemented the evidence with data from recently published RCTs.

Type of participants (Population of interest)

Suitable studies included men and women aged 18 and above diagnosed with MS. Eligible MS diagnosis was RRMS. CIS patients were not included in this report; however, Appendix 3 lists identified studies that included CIS patients. We excluded studies with patients with primary progressive MS and radiologically isolated syndrome. Studies that included both eligible patients, and patients from our exclusion criteria were included if results were presented separately for each type of patients (so that we could extract results for patients who fitted our inclusion criteria).

Types of interventions

The following medicines were the interventions of interest: dimethyl fumarate, teriflunomide, interferon beta, peg-interferon, glatiramer acetate, natalizumab, fingolimod, and alemtuzumab.

Comparisons

Eligible comparison groups were either placebo or one of the medicines listed above.

Types of outcome measures

The outcomes of interest were:

Primary outcomes

- Clinical relapses
- Disability progression measured using the EDSS
- Mortality
- Serious adverse events

Secondary outcomes:

- Withdrawal from study due to adverse events
- Stay at hospitals
- Health related quality of life measured with EQ-5D

Literature search

The research librarian (in collaboration with the project team) conducted a peer-reviewed literature search using index terms (Medical Subject Headings and EMTREE terms) and free text terms relating to the population and the interventions of interest. The last date of the literature search was 9/11/2015. Full literature search strategies are presented in Appendix 1. We did not use any language restrictions in the literature search.

We searched the following databases:

- Ovid MEDLINE(R) In-Process & Other Non-Indexed Citations and Ovid MEDLINE(R)
- Embase
- Cochrane Library; Cochrane Database of Systematic Reviews, Other Reviews, Technology Assessments, Cochrane Central Register of Controlled Trials (Central)
- Centre for Reviews and Dissemination; DARE, HTA
- ISI web of Science
- PubMed (epub ahead of print)
- Epistemonikos

We searched also the following websites:

- Canadian Agency for Drugs and Technologies in Health (CADTH)
- Agency for Healthcare Research and Quality (AHRQ),
- FinOHTA- Finnish Office for Health Technology Assessment
- Statens beredning för medicinsk utvärdering (SBU)
- EUnetHTA POP database (POP = Planned and Ongoing Projects)
- PROSPERO Centre for Reviews and Dissemination

We checked bibliographies of selected articles for additional publications meeting our inclusion criteria. Finally, we searched the WHO ICTRP and ClinicalTrials.gov to identify relevant ongoing or unpublished trials.

We contacted the companies with marketing authorization in Norway for the MS medicines included in order to get additional information.

Selection and assessment of publications

Selection of publications

Unless stated otherwise, two persons independently carried out the selection processes.

Selection of HTA or SR reports

Two persons read titles and abstracts retrieved by the literature search, and excluded obviously irrelevant literature. Based on information provided in abstracts, one person organised the publications depending on how many medicines were apparently examined. Abstracts looking at two, three, or more than three drugs were grouped together. If we lacked information in the abstract to know which medicine were assessed, articles were classified in the "several drugs category" (more than three drugs). One person sorted all abstracts in the "several drugs category" according to the date of publication from the newest to the oldest. Two persons read full-text articles of the "several drugs category" by publication chronological order (from newest to oldest). Hence, we were able to include the most recently published HTA report that met all our inclusion criteria.

Selections of RCT publications

Two persons examined all titles and abstracts retrieved by the literature search for possibly relevant RCTs published after the selected HTA, and excluded obviously irrelevant titles and abstracts. Two persons read full-text articles of selected publications. We included articles that met our inclusion criteria. The same process was used to select publications sent by companies having market authorization for MS medicines in Norway.

Throughout the selection process, any disagreement was discussed to reach an agreement.

Assessment of included publications

Quality assessment of selected HTA

We assessed the quality of the SR part of the identified HTA using the checklist for SR in the handbook of The Norwegian Knowledge Centre for the Health Services (21).

Risk of bias of RCTs

We did not perform risk of bias assessments for the RCTs included in the selected high quality HTA report. Instead, we report the risk of bias assessments conducted by the HTA authors. The domains of risk of bias assessed in the HTA report were similar to the Cochrane Collaboration tool for assessing risk of bias (22) (randomization, allocation concealment, double-blinding, baseline characteristics similarity, outcome measures, withdrawals, use of intention-to-treat analysis, and source of funding). For the newer RCTs that we supplemented, we used the Norwegian Knowledge Centre for the Health Services tool to assess risk of bias (23). That tool is based on Cochrane risk of bias tool (22).

The assessment of risk of bias of included RCTs was carried out by one person and checked by another. For the evaluation of risk of bias provided by the HTA report, one author extracted the assessment data, and another verified the data. Any disagreements were discussed to reach consensus.

Data collection and analysis

Data extraction

One person extracted predefined data from the selected publications, and a second checked the data extraction for accuracy.

Data extraction from HTA/SR

We extracted the following data from the selected HTA report: publication information (authors, publication details), date of the literature search, characteristics of included studies (study design, origin, setting, comparisons and endpoints investigated, follow-up range of included studies), and information on quality assessment.

Data extraction from RCTs

We extracted the following data from included RCTs: information on publication (authors , publication details); RCT description (clinical trial identification, design and setting, source of funding); participants characteristics (age and gender, MS diagnosis, inclusion and exclusion criteria, and baseline characteristics); description of intervention and comparison groups (numbers of participants in each group, doses, administration method); and outcomes (primary and secondary endpoints assessed, definitions used, length of follow-up, measurements of outcomes such as number of events, means, corresponding standard deviations).

For RCTs included in the HTA, for each individual RCT, we extracted the data reported in the HTA publication. To assess accuracy, one person compared the information given by the HTA report with the original study publication of seven randomly chosen RCTs. All the data presented in the HTA were identical to the original publications. For RCTs identified after the HTA literature search, we extracted the data from the primary publications.

Statistical analyses and presentation of results

Measures of treatment effect

We expressed the comparative effectiveness of the treatments as the relative risk (RR) for dichotomous outcomes, annualised rate ratios (ARR) for count data and the mean difference (MD) for continuous outcomes. For all outcomes 95% confidence intervals (CI) or credible intervals (CI) were calculated for the RR, ARR, MD. The credible interval is the Bayesian analogue to confidence intervals used in traditional frequentist statistical approaches. We considered a difference to be "significant" if the CrI did not include RR =1 or MD=0.

For count data (number of relapses), we used a Poisson regression based approach to obtain the annualised rate ratios (ARR) from the total number of relapses and patient-years of follow-up.

Dealing with missing data

For the endpoint "number of relapses" we performed imputations to derive needed values where included trials did not report the total number of relapses or exposure time (person-years). Missing number of total relapses were derived using the exposure time (person-years) and the reported mean ARR values. For missing exposure-time (in person-years), the values were imputed using treatment duration and number of patients completing the study (100% was assumed in cases where the percentage of completers was not reported).

For disability progression, measured as a dichotomous outcome, we assumed that participants who dropped out experienced the event (a likely scenario). For all other

endpoints, we did not perform imputations for missing data. We based the statistical analyses on the intention to treat principle (all participants analysed in the group to which they were allocated, and all available data included in the analyses).

The statistical analysis was based on binomial likelihoods (dichotomous outcomes), poisson likelihoods (count outcomes), and normal likelihood (continuous outcomes), with vague priors for the trial baselines, basic parameters (normal distribution with mean 0 and standard deviation 0.0001) and the random effects standard deviation (uniformly distributed in the interval 0 to 2), and takes the correlation structure induced by multi-arm trials into account. We used a random effects model. We checked for incoherence between direct and indirect evidence by "node-splitting" (24). We calculated the direct and indirect estimates of effect and the corresponding Bayesian "P-values" for incoherence.

We ranked the different treatments in terms of their likelihood of leading to the best results for each primary endpoint. We based the rankings on the surface under the cumulative ranking curve (SUCRA) (25). We interpreted the rankings cautiously taking into account the quality of evidence.

We performed sensitivity analyses where participants who dropped out were excluded from the analyses of the sustained disability progression, to base the analyses only on the available data.

Data synthesis

First, we conducted pairwise meta-analyses for each available outcome and, for each identified intervention vs. control group comparison. This was done using a traditional frequentist statistical approach assuming random effects models using the software RevMan 5.3. Hereafter, we refer to this method as the "pairwise comparisons method". Further, we combined direct and indirect evidence, and performed a network-meta-analysis (19). For that, we used a Bayesian method based on Markov Chain Monte Carlo simulation. This method is, hereafter, referred to as the "network meta-analysis approach". This was done using Winbugs version 1.4.3 (Imperial College and MRC, UK).

Grading the quality of evidence

Two review authors assessed independently the quality of the evidence for each selected outcome. We used Grading of recommendations Assessment, Development, and Evaluation (GRADE) to assess the quality of the direct evidence, indirect evidence, and the combined evidence from the NMA (26).

First, we graded the evidence for all comparisons with available direct evidence. Then, we graded the comparisons for which we had indirect evidence. To grade the indirect evidence, we considered the direct evidence that contributed to that indirect evidence. For example, the indirect evidence comparing a medicine A with a medicine C might have been obtained with direct evidence comparing medicines A and B, and B with C. The grade of the indirect evidence for the comparison A and C was based on the grade of the direct evidence on A and B, and B and C. The grade of the indirect evidence on A versus C was the lowest grade of all the direct evidence that contributed to that comparison.

To select the direct evidence that might have contributed to the indirect evidence, we chose the evidence that involved fewest head-to-head comparisons. For example, for indirect evidence comparing A to C, one might also have evidence comparing A to D, D to E and E to C. This example involves three head-to-head comparisons compared to the two presented above (A with B, and B with C). The indirect evidence with fewer head-to-head comparisons is referred to as first order loops. If more than one first order loops were available, we chose the loop with the lowest available quality. This was a conservative approach.

For a specified comparison, the grade of the network meta-analysis evidence was the highest GRADE between the direct and indirect evidence for that comparison.

Due to time constraint, we graded the quality of the evidence only for annual relapse rate, disability progression (when examining disability progression as a dichotomous variable: considering whether someone had been less disabled or not when using a certain treatment) and withdrawal due to adverse events. The first two outcomes were the two outcomes used in the economic evaluation. Withdrawal due to adverse events is also an important outcome as it measures the risk of adverse event(s) outweighing the benefit of the treatment to the point of causing withdrawal from treatment.

GRADE provides specific criteria to consider when rating the quality of evidence. This includes the strength of the study design, possible risk of bias, imprecision and inconsistency of the estimates, and indirectness and magnitude of effect, dose response gradient and potential confounding factors. The overall quality of the evidence was classified as high, moderate, low, or very low for each outcome. The definition for each category is described in the following table.

Table 2. Definition of each category for GRADE

Grade	Definition
High	We are very confident that the true effect lies close to that of the estimate of effect
Moderate	We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of effect, but there is a possibility that it is substantially different
Low	Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect
Very low	We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Clinical evaluation - Results

Result of literature search

Results of the search and selection process

We selected the evidence for this report in two stages, first identifying relevant SRs or HTA reports (Figure 1), and then supplementing the evidence of the identified HTA with more up to date information (Figure 2).

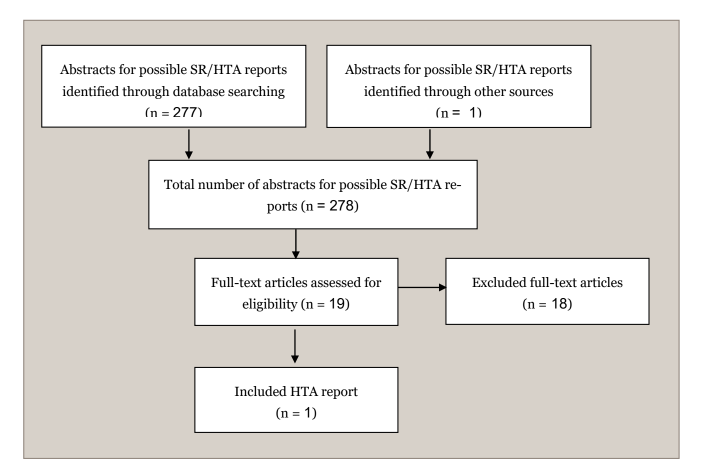


Figure 1. Flow diagram for the selection of possible systematic reviews (SR) or health technology assessment (HTA) reports

When looking for possible SR or HTA reports, the literature search retrieved 277 records, and we found one extra record. After abstract selection, and assessing 19 full-text articles, we included one HTA. This was a recent HTA report (literature search carried out in October 2013). To supplement the HTA's information with more up to date evidence, we searched for additional RCTs published from 2013 to the last date of our literature search (9/11/2015).

The literature search for RCTs identified 644 records. We supplemented this search with two records identified in reference lists, and one RCT provided by a pharmaceutical company. After the selection process, we included fifteen publications on eleven RCTs.

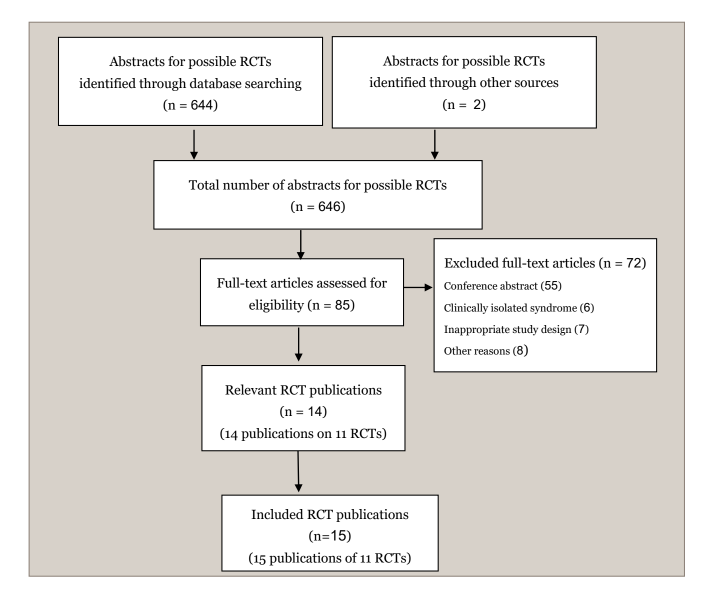


Figure 2. Flow diagram for the selection of possible randomised clinical trials (*RCT*) published after the included health technology assessment report

Included studies

The included health technology assessment report

Some of the evidence presented in this report was extracted from a previously published HTA report (27). This publication is described in Table 3. It summarised evidence from RCTs assessing mono- and combination therapies of MS-medicines. We included data from 26 RCTs (only the RCTs that examined MS monotherapies). The participants were RRMS patients, with a mean age ranging between 29 and 41 years old. They were followed for a period ranging from 16 weeks to 3.5 years, and were in majority women.

Date of literature search	October 2013
Study types included	RCTs (Number of included monotherapy RCTs: 26)
Participants	 All studies included patients with RRMS. One study included patients with clinically isolated syndrome (CIS), one study included patients with progressive-relapsing MS (PRMS), one study included patients with secondary-progressive, and one study included patients with second-ary-progressive MS and progressive-relapsing MS. Randomized sample size: 75 to 1430. Female participants: 64% to 84% Mean age: 29 to 41 years
Intervention (number of	Alemtuzumab (three)
unique RCTs)	Dimethyl fumarate (two) Fingolimod (three)
_	Glatiramer acetate (eight)
	Interferon beta-1a subcutaneous (nine)
	Interferon beta-1a intramuscular (nine)
	Interferon beta-1b (five)
	Natalizumab (one)
	Teriflunomide (two)
Comparison	Placebo
	One of the drugs listed above
Outcome	- Relapse - Disability progression
	- MRI lesions
	- Adverse events
	- Serious adverse events
	- Withdrawal due to adverse events
	- Quality of life
Follow-up	16 weeks to 3.5 years.
Quality assessment	This publication was assessed to be of high quality

Table 3. Characteristics of the included HTA report

The included primary studies

We present an overview of RCTs that constitute our evidence base in Table 4. Further details on both the primary studies included in the above-mentioned HTA report, and those we identified are provided in Appendix 2.

Altogether, we included 37 studies; 26 from the selected HTA report (27), and 11 RCTs from our supplementary search. All RCTs included RRMS patients. Treatment histories varied, with 11 RCTs confined to treatment-naive patients, 4 included treatment experienced participants, 11 combined treatment naïve and treatment experienced patients, and treatment history was unclear in 9 studies. We had information for 39 comparisons including active treatments versus placebo, and active treatments compared with each other.

Many of the published studies did not examine medications separating first- and second- line treatments. Studies compared first-line treatments and second-line treatments (28-32). Other studies examined first-line treatments in patients who had taken other medications before (33-36). Two studies investigated second-line treatments in a population that comprised treatment naive patients (i.e. patients who had not received a first-line treatment) (37, 38).

Excluded studies

Excluded studies and reasons for exclusion are presented in Appendix 3.

Ongoing studies and other relevant literature

We searched the WHO ICTRP and ClinicalTrials.gov to identify relevant ongoing or unpublished trials. The result of this search is presented in Appendix 4.

Table 4. Characteristics of included randomised clinical trials

Name (publication) (reference)			
Study design	ised)	history	
CAMMS223 (2008)(28)	- Alemtuzumab 12 mg IV q.d., 5 consecutive days at 1st month, 3	Treatment-	3 years
Rater-blinded, in 49 centres in Europe	consecutive days at months 12 and 24 $(n = 113)$	naive	
and US	- Alemtuzumab 24 mg IV q.d. (n = 110)		
	- Interferon beta-1a 44 mcg SC t.i.w. $(n = 111)$		
CARE-MS I(2012) (29)	- Alemtuzumab 12 mg IV q.d., 5 consecutive days at month 0, 3	Treatment-	2 years
A rater-blinded, in 101 centres in 16	consecutive days at month 12 ($n = 386$)	naive	
countries including Europe, Canada,	- Interferon beta-1a 44 mcg SC t.i.w. $(n = 195)$		
and US.			
CARE MS II (2008)(28)	- Alemtuzumab 12 mg IV q.d., 5 consecutive days at month 0, 3	Treatment-	2 years
Rater-blinded, in 194 academic medi-	consecutive days at month 12 (n=436)	experienced	
cal centres and clinical practices in 23	Alemtuzumab 24 mg IV q.d., 5 consecutive days at month 0, 3		
countries including Europe, Canada,	consecutive days at month 12 (n=173)		
and US.	- Interferon beta-1a 44 mcg SC t.i.w. (n=231)	! .	
DEFINE (2012) (33)	- Dimethyl fumarate 240 mg oral twice daily (n = 410) [total 480	Mixed	2 years
Double-blind, in 28 countries includ-	mg/day]		
ing Europe, Canada, and US	- Dimethyl fumarate 240 mg oral three times daily (n = 416) [total		
	720 mg/day]		
	- Placebo (n = 408)		
CONFIRM (2012) (34)	- Dimethyl fumarate 240 mg b.i.d, (n=359)	Mixed	2 years
Rater-blinded, in 200 research sites in	- Dimethyl fumarate 240 mg three times daily (n=345), subcuta-		
28 countries including Europe and	neous daily injections of 20 mg of glatiramer acetate for 96 weeks		
North America	(n=350)		
	- Placebo (n=363)		

Name (publication) (reference) Study design	Intervention versus comparison (n=number random- ised)	Treatment history	Follow-up
FREEDOMS (2010) (37) Double-blind, multi-centre in Aus- tralia, Canada, Europe, and South Af- rica (138 centers in 22 countries)	- Fingolimod oral 0.5 mg q.d. (n = 425) - Fingolimod oral 1,25 mg q.d. (n = 429) - Placebo (n = 418)	Mixed	2 years
TRANSFORMS (2010) (38) Double-blind, in 172 centres in 18 countries including Canada, Australia, Europe, and US.	 Fingolimod oral 0.5 mg q.d. (n=431) Fingolimod oral 1.25 mg q.d. (n=426) Interferon beta-1a 30 mcg IM q.w. (n=435) 	Mixed	1 year
<i>Saida et al. (2012) (39)</i> Double-blind, multicentre in Japan	 Fingolimod oral 0.5 mg q.d. (n=57) Fingolimod oral 1.25 mg q.d. (n=57) Placebo (n=57) 	Unclear	6 months
FREEDOMS II (2014) (40, 41) Double-blind, in 117 academic and ter- tiary referral centres in 8 countries, most patients included in the USA	 Fingolimod 0.5 mg oral q.d. (n=358) Fingolimod 1.25 mg oral q.d. (n=370) Placebo (n=355) 	Unclear	2 years
Johnson et al. (1995) (42) Double-blind, in 11 centres in the US	- Glatiramer acetate 20 mg SC q.d (n =125) - Placebo (n=126)	Treatment- naive	2 years
<i>Comi et al. (2001)(43)</i> Double-blind, in 7 countries	- Glatiramer acetate 20 mg SC q.d. (n=119) - Placebo (n=120)	Unclear	9 months
REGARD (2008) (44) Open-label, rater-masked. 81 centres in 14 countries including Canada, South America, and Europe	- Glatiramer acetate 20 mg SC q.d. (n=378) - Interferon beta-1a 44 mcg SC t.i.w. (n=386)	Treatment- naive	96 weeks
BECOME (2009) (45) Rater-blinded, in one centre in the US	- Glatiramer acetate 20 mg SC q.d. (n = 39) - Interferon beta-1b 250 mcg SC every other day (n = 36)	Treatment- naive	2 years

Name (publication) (reference) Study design	Intervention versus comparison (n=number random- ised)	Treatment history	Follow-up
BEYOND (2009)(46)	- Glatiramer acetate 20 mg SC q.d. (n = 448)	Treatment-	2 to 3,5
A rater-blinded, in 198 centres in 26 countries worldwide.	 Interferon beta-1b 250 mcg SC every other day (n = 897) Interferon beta-1b 500 mcg SC every other day (n = 899) 	naive	years
<i>Calabrese et al. (2012)(47)</i> Rater-blinded, single-centre in Italy	- Glatiramer acetate 20 mg SC q.d. (n = 55) - Interferon beta-1a 44 mcg SC t.i.w. (n = 55) - Interferon beta-1a 30 mcg IM q.w. (n = 55)	Unclear	2 years
<i>GALA (2013)(35)</i> Double-blind study, in 142 sites in 17 countries	- Glatiramer acetate sc 40mg (1ml) tiw (n=943) - Placebo (n=461)	Mixed	1 year
<i>CombiRx (2013) (48)</i> Double-blind, in 68 sites, both private practice and academic, in the USA and Canada	 Interferon beta-1a 30µg IM q.d and glatiramer acetate (GA) 20mg q.d (n=499) (not considered)) Glatiramer acetate 20mg q.d (n=259) Interferon beta-1a 30µg IM q.w (n=250) These interventions were compared one with another 	Treatment- naïve	3 years
<i>MSCRG (1996)(49)</i> Double-blind, in 4 centres in the US	- Interferon beta-l a 30 mcg IM q.w. (n=158) - Placebo (n=143)	Treatment- naive	2 years
EVIDENCE (2002) (50) Rater-blinded, in 56 centres in Europe, Canada, and US.	- Interferon beta-1a 30 mcg IM q.w. (n = 338) - Interferon beta-1a 44 mcg SC t.i.w. (n = 339)	Unclear	24 weeks
INCOMIN (2002) (51) Open label, rater-masked, in 15 cen- tres in Italy	 Interferon beta-1a 30 mcg IM q.w. (n = 92) Interferon beta-1b 250 mcg SC every other day (n = 96) 	Treatment- naive	2 years
<i>Clanet et al.</i> (2002) (52) Double-blind, dose-comparison study. In 38 centers in Europe	 Interferon beta-1a 30 mcg IM once weekly (n=402) Interferon beta-1a 60 mcg IM once weekly N=(400) 	Unclear	At least 3 years

Name (publication) (reference) Study design	Intervention versus comparison (n=number random- ised)	Treatment history	Follow-up
<i>Kappos et al. (2011) (36)</i> 79 centres in 20 countries in North America, east-central Europe, Asia, western Europe, and Latin America.	 Ocrelizumab 600 mg IV day 1 and 15 (n=55, not our scope) Ocrelizumab 600 mg IV day 1 and 15 (n=55, not our scope) Interferon beta-1a 30 mcg IM q.d. (n=55) Placebo (n=54) 	Mixed	24 weeks
<i>Mokhber et al. (2013) (53)</i> <i>S</i> ingle center in Iran	 Interferon beta-1a (Avonex) 30 mcg once per week IM injection; (n=23) Interferon beta-1a (Rebif) 44 mcg t.i.w. SC injection; (n=23) Interferon beta-1a (Betaferon) 0.25 mg every other day SC injection (n=23) 	Treatment- naive	1 year
BRAVO (2014) (54) In 18 countries	 Laquinimod 0.6 mg capsule q.d. (n=434)[not our scope] Interferon beta-1a IM 30 mcg once-weekly injection (n = 447) Placebo (matching laquinimod) (n = 450) 	Mixed	2 years
PRISMS (1998) (55) Double-blind, in 22 centres in 9 coun- tries including Australia, Canada, and Europe	 Interferon beta-1a 22 mcg SC t.i.w.(n=189) Interferon beta-1a 44 mcg SC t.i.w. (n=184) Placebo (n=187) 	Treatment- naive	2 years
<i>IMPROVE (2010) (56)</i> Double-blind, multi-centre, multi- country in European countries.	- Interferon beta-1a 44 mcg SC t.i.w. (n = 120) - Placebo (n = 60)	Unclear	16 weeks
<i>IFNB-MS (1993) (57)</i> Multi-centre Canada and the US.	 Interferon beta-1b 250 mcg SC every other day (n = 124) Interferon beta-1b 50 mcg SC every other day (n=125) Placebo (n = 123) 	Treatment- naïve	3 years
<i>Etemadifar et al. (2006)(58)</i> Rater-blinded, neurology outpatient clinics in Iran	 Interferon beta-1b 250 mcg SC every other day (n = 30) Interferon beta-1a 30 mcg IM q.w. (n = 30) Interferon beta-1a 44 mcg SC t.i.w. (n = 30) 	Unclear	2 years
ADVANCE study(2014) (59) Double-blind, in 26 countries, in north/south America, Europe, India	 Peg-interferon beta-1a 125 mcg SC once every 2 weeks (n=512) Peg-interferon beta-1a 125 mcg SC once every 4 weeks (n=500) Placebo (n=500) 	Mixed	2 years

 $mg = milligrams, mcg = micrograms, SC = subcutaneous; \ q.d. = once \ daily, \ q.w. = . \ once \ weekly, \ t.i.w. = three \ times \ weekly, \ IM = intra \ muscular \ muscular$

Name (publication) (reference) Study design	Intervention versus comparison (n=number random- ised)	Treatment history	Follow-up
AFFIRM (2006) (60) Double-blind, in 99 centres in Europe, North America, Australia, and New Zealand.	- Natalizumab 300 mg IV every 4 weeks (n = 627) - Placebo (n = 315)	Unclear	2 years
<i>Gobbi et al (2013) (31)</i> Rater blinded. One centre, Switzer- land.	 Continue on natalizumab 300 mg IV q.m. (n=10) Switch to interferon beta-1b 250 mcg every other day (n=9) 	Treatment experienced	1 year
RESTORE (2014) (61) Randomized partially, in North Amer- ica and Europe	 Natalizumab 300 mg IV every 4 weeks (n=45) Alternate immunomodulatory therapy (n=88) (not our scope) Placebo IV every 4 weeks (n=42) 	Treatment experienced	24 weeks
Zecca et al. (2014) (32) Rater-blinded, parallel-group study, single center, Switzerland	 Continue Natalizumab monthly intravenous (i.v.) 300 mg (n=10) De-escalate to interferon beta-1b subcutaneous (s.c.) 250 mcg every other day (n=9) 	Treatment experienced	1 year
O'Connor et al (2006) (62) Double-blind. Centres in Canada	- Teriflunomide oral 7 mg q.d.(n=61) - Teriflunomide oral 14 mg q.d.(n=57 - Placebo (n=61)	Treatment- naive	36 weeks
TEMSO (2011) (63, 64) Double-blind, in 127 centres in 21 countries including Canada, Europe, and US.	- Teriflunomide oral 7 mg q.d. (n=365) - Teriflunomide oral 14 mg q.d. (n=358) - Placebo (n=363)	Mixed	108 weeks
<i>TOWER (2014) (65)</i> Double-blind, in 189 centres mainly hospital-based sites in 26 countries	- Teriflunomide 14 mg once daily (n=372) - Teriflunomide 7 mg once daily (n=408) - Placebo once daily (n=389)	Mixed	Up to 48 weeks
TENERE (2014) (66) Rater-blinded study, multicentre study	 Teriflunomide 14 mg oral once daily (n=111) Teriflunomide 7 mg oral once daily (n=109) Interferon beta-1a 44mcg s.c three times/week (n=104) 	Mixed	Up to 48 weeks

Effects of intervention(s)

We describe here the effects of the examined MS disease modifying medicines on outcomes.

The GRADE evaluation is described in detail in Appendix 5. Results of the full network meta-analysis for all possible comparisons for all outcomes are given in Appendix 6.

Annualised relapse rate

We present here the results obtained using the "network meta-analysis approach" (Bayesian method). We found similar results using the "pairwise comparison method" (Frequentist approach). Those results are presented in Appendix 7.

Figure 3 shows the available network of evidence for annualised relapse rate. The thickness of the line is proportional to the amount of evidence for that comparison. In total, 19 MS treatment strategies and placebo were examined.

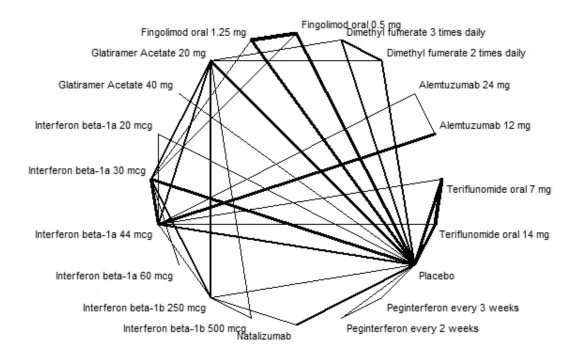


Figure 3. Evidence network for annualised relapse rate

Active treatments versus placebo

Fifteen treatments were compared to placebo (Table 5). Results from direct, and indirect evidence, and from the whole network are consistent (except for teriflunomide oral 7 mg). All active treatments examined were more effective than placebo against relapse. The highest effect against annual relapse was seen for alemtuzumab 12 mg IV q.d. When considering results we had high quality evidence for, the relative risk for annual relapse ranged between 0.29 (95% CI: 0.23; 0.35) for alemtuzumab 12 mg IV q.d, and 0.86 (0.7 to 1.06) for interferon beta-1a 60 mcg IM q.w, compared to placebo.

	Direct evid	Direct evidence Indirect evidence Network met		Indirect evidence		analysis
Interventions	RR (95% CI)	GRADE	RR (95% CI)	GRADE	RR (95% CI)	GRADE
Interferon beta-1a 22 mcg	0.69 (0.57 to 0.83)	Moderate	NA	NA	0.69 (0.57 to 0.83)	Moderate
Interferon beta-1a 30 mcg	0.76 (0.65 to 0.89)	High	0.87 (0.75 to 1.01)	Moderate	0.82 (0.73 to 0.91)	High
Interferon beta-1a 44 mcg	0.67 (0.54 to 0.80)	High	0.61 (0.52 to 0.72)	Very low	0.64 (0.56 to 0.72)	High
Glatiramer acetate 20mg	0.70 (0.60 to 0.82)	High	0.60 (0.52 to 0.70)	Moderate	0.65 (0.59 to 0.73)	High
Glatiramer acetate 40mg	0.66 (0.52 to 0.82)	High	NA	NA	0.66 (0.52 to 0.82)	High
Dimethyl fumarate 240 mg 2.i.d	0.5 (0.42 to 0.6)	High	NA	NA	0.5 (0.42 to 0.6)	High
Dimethyl fumarate 240 mg t.i.d	0.5 (0.42 to 0.6)	High	NA	NA	0.5 (0.42 to 0.6)	High
Teriflunomide oral 7 mg	0.73 (0.64 to 0.84)	High	1.12 (0.78 to 1.57)	Moderate	0.77 (0.68 to 0.9)	High
Teriflunomide oral 14 mg	0.67 (0.58 to 0.78)	High	0.57 (0.39 to 0.83)	Low	0.67 (0.58 to 0.77)	High
Fingolimod oral 0.5 mg	0.49 (0.41 to 0.57)	High	0.38 (0.27 to 0.51)	Moderate	0.46 (0.39 to 0.54)	High
Fingolimod oral 1.25 mg	0.43 (0.37 to 0.51)	High	0.53 (0.39 to 0.84)	Moderate	0.45 (0.39 to 0.53)	High
Peg-interferon beta-1a 125 mcg 1/ 2 w	0.65 (0.49 to 0.85)	High	NA	NA	0.65 (0.49 to 0.85)	High
Peg-interferon beta-1a 125 mcg 1/4 w	0.73 (0.56 to 0.95)	High	NA	NA	0.73 (0.56 to 0.95)	High
Natalizumab	0.30 (0.25 to 0.36)	Moderate	0.0002 (0.00 to 0.07)	Very low	0.3 (0.24 to 0.36)	Moderate
Interferon beta-1b 250 mcg	0.65 (0.51 to 0.83)	Moderate	0.67 (0.55 to 0.79)	Very low	0.66 (0.57 to 0.76)	Moderate
Alemtuzumab 24 mg IV q.d.	NA	NA	0.16 (0.1 to 0.25)	Low	0.16 (0.1 to 0.25)	Low
Alemtuzumab 12 mg IV q.d	NA	NA	0.29 (0.23 to 0.35)	High	0.29 (0.23 to 0.35)	High
Interferon beta-1b 500mcg SC 1/2 d	NA	NA	0.62 (0.51 to 0.74)	Moderate	0.62 (0.51 to 0.74)	Moderate
Interferon beta-1a 60 mcg IM q.w.	NA	NA	0.86 (0.7 to 1.06)	High	0.86 (0.7 to 1.06)	High

RR= relative ratio, *CI*= confidence interval, *mg*= milligrams, *mcg*= micrograms, *SC*= subcutaneous, *IM*= intra muscular, *q.d.*= once daily, *q.w.*=once weekly, *t.i.w.*= three times weekly, *2.i.d*= two times daily, *t.i.d*= three times daily, *1/2* w=once every 2 weeks, *1/4* w=once every 4 weeks, *1/2* d= once every two days, *NA*=Not applicable (no available data).

Active treatments compared with each other

We had information on 24 head-to-head comparisons of active treatments (Table 6). Most results (except interferon beta-1a 44mcg versus alemtuzumab 24 mg and interferon beta-1a 22 mcg; and for teriflunomide oral 7 mg versus interferon beta-1a 44 mcg) were similar for direct, and indirect evidence, and for the whole network. When considering statistically significant results for which we had high quality of evidence, we found that some treatments were more effective than others against relapses: interferon beta-1a 44 mcg was less effective than alemtuzumab 12 mg (RR; 95% CI= 2.21; 1.90 to 2.64). Fingolimod oral 0.5 mg and fingolimod oral 1.25 mg performed better than interferon beta-1a 30 mcg, with RRs (95% CI) of 0.57 (0.47 to 0.67) and 0.55 (0.47 to 0.66), respectively. Furthermore, dimethyl fumarate 240 mg two times and three times daily were more effective than glatiramer acetate 20mg, with RRs of 0.77 (0.63 to 0.93) and 0.77 (0.64 to 0.93), respectively.

		Direct evidence		Indirect evidence		Network meta-analysis	
Intervention	Comparison	RR (95% CI)	GRADE	RR (95% CI)	GRADE	RR (95% CI)	GRADE
Alemtuzumab 24 mg	Alemtuzumab 12 mg	0.55 (0.35 to 0.86)	Low	NA	NA	0.55 (0.35 to 0.86)	Low
Interferon beta-1a 44 mcg	Alemtuzumab 12 mg	2.21 (1.9 to 2.64)	High	NA	Low	2.21 (1.9 to 2.64)	High
Interferon beta-1a 44 mcg	Interferon beta-1a 22 mcg	0.68 (0.56 to 0.83)	Moderate	0.43 (0.33 to 0.55)	Moderate	0.92 (0.76 to 1.11)	Moderate
Interferon beta-1a 44 mcg	Interferon beta-1a 30 mcg	0.76 (0.63 to 0.93)	High	0.79 (0.65 to 0.95)	Very low	0.78 (0.68 to 0.89)	High
Interferon beta-1a 60 mcg	Interferon beta-1a 30 mcg	1.05 (0.88 to 1.25)	Moderate	NA	NA	1.05 (0.88 to 1.25)	Moderate
Glatiramer acetate 20mg	Interferon beta-1a 30 mcg	0.79 (0.61 to 1.02)	Moderate	0.80 (0.69 to 0.93)	Moderate	0.8 (0.7 to 0.91)	Moderate
Fingolimod oral 0.5 mg	Interferon beta-1a 30 mcg	0.48 (0.35 to 0.64)	High	0.60 (0.50 to 0.73)	Moderate	0.57 (0.47 to 0.67)	High
Fingolimod oral 1.25 mg	Interferon beta-1a 30 mcg	0.63 (0.46 to 0.90)	High	0.52 (0.43 to 0.63)	Moderate	0.55 (0.47 to 0.66)	High
Interferon beta-1b 250 mcg	Interferon beta-1a 30 mcg	0.71 (0.53 to 0.91)	Moderate	0.85 (0.71 to 1.03)	Very low	0.81 (0.69 to 0.93)	Moderate
Glatiramer acetate 20mg	Interferon beta-1a 44 mcg	1.02 (0.83 to 1.28)	Moderate	0.98 (0.82 to 1.18)	Very low	1.02 (0.9 to 1.18)	Moderate
Teriflunomide oral 7 mg	Interferon beta-1a 44 mcg	1.72 (1.24 to 2.44)	Moderate	1.13 (0.93 to 1.34)	Low	1.21 (1.02 to 1.47)	Moderate
Teriflunomide oral 14 mg	Interferon beta-1a 44 mcg	0.91 (0.62 to 1.36)	Low	1.06 (0.89 to 1.31)	Moderate	1.04 (0.87 to 1.27)	Moderate
Interferon beta-1b 250 mcg	Interferon beta-1a 44 mcg	0.81 (0.46 to 1.43)	Very low	1.00 (0.83 to 1.18)	Moderate	1.03 (0.88 to 1.22)	Moderate
Ddimethyl fumarate 240 mg 2.i.d	Glatiramer acetate 20mg	0.59 (0.38 to 0.90)	High	0.63 (0.40 to 0.98)	Moderate	0.77 (0.63 to 0.93)	High
Dimethyl fumarate 240 mg t.i.d	Glatiramer acetate 20mg	0.53 (0.35 to 0.79)	High	0.78 (0.50 to 1.25)	Moderate	0.77 (0.64 to 0.93)	High
Interferon beta-1b 250 mcg	Glatiramer acetate 20mg	1.07 (0.90 to 1.27)	Moderate	0.92 (0.75 to 1.14)	Very low	1.01 (0.88 to 1.16)	Moderate
Interferon beta-1b 500 mcg	Glatiramer acetate 20mg	0.95 (0.8 to 1.12)	Moderate	NA	NA	0.95 (0.8 to 1.12)	Moderate
Dimethyl fumarate 240 mg t.i.d	Dimethyl fumarate 240 mg 2.i.d	1.01 (0.82 to 1.23)	Moderate	NA	NA	1.01 (0.82 to 1.23)	Moderate
Teriflunomide oral 14 mg	Teriflunomide oral 7 mg	0.86 (0.74 to 1.)	Moderate	NA	NA	0.86 (0.74 to 1.)	Moderate
Fingolimod oral 1.25 mg	Fingolimod oral 0.5 mg	0.98 (0.83 to 1.17)	Moderate	NA	NA	0.98 (0.83 to 1.17)	Moderate
Peginterferon beta-1a 125 mcg 1/4 w	Peginterferon beta-1a 125 mcg 1/2 w	1.13 (0.84 to 1.52)	Moderate	NA	NA	1.13 (0.84 to 1.52)	Moderate
Interferon beta-1b 250 mcg	Natalizumab	NE	Very low	2.17 (1.71 to 2.76)	Moderate	2.22 (1.76 to 2.81)	Moderate
Interferon beta-1b 500 mcg	Interferon beta-1b 250 mcg	0.93 (0.8 to 1.1)	Moderate	NA	NA	0.93 (0.8 to 1.1)	Moderate

Table 6. Relative risk for annual relapse for active MS treatments compared to others for comparisons with available direct evidence

RR= relative ratio, CI= confidence interval, mg= milligrams, mcg= micrograms, SC= subcutaneous, IM= intra muscular, q.d.= once daily, q.w.=once weekly, t.i.w.= three times weekly, 2.i.d= two times daily, t.i.d= three times daily, 1/2 w=once every 2 weeks, 1/4 w=once every 4 weeks.

NE= *Not estimable (Estimate of difference for direct evidence is not estimable due to 0 events in the Natalizumab group)*

Disability progression

We examined, first, disability progression as a dichotomous variable, considering whether someone had been less disabled or not when using a certain treatment. The results obtained using the "network meta-analysis approach" are presented here. These results are consistent with results found with the "pairwise comparison method". The "pairwise comparison method" results are presented in Appendix 7.

The network of evidence available for disability progression is presented in Figure 4. We had evidence for 18 treatment strategies and placebo.

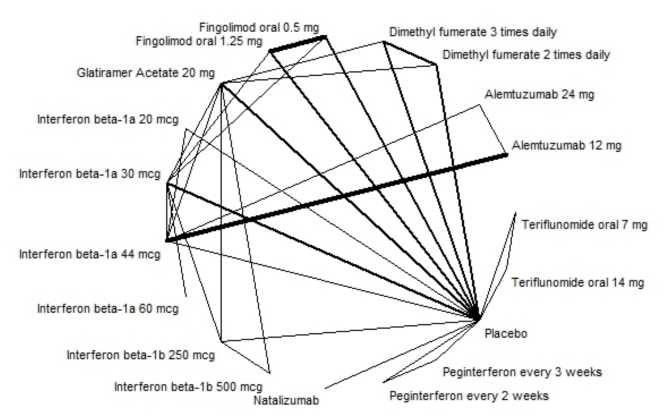


Figure 4. Evidence network for disability progression

Active treatments versus placebo

Table 7 compares results obtained when considering direct, indirect evidence and the whole network. It shows that results were similar. Seventeen treatments were compared to placebo. For four of these, we had high quality evidence, and they were all more effective than placebo against disability progression. The network meta-analysis RRs for disability progression were 0.65 (95% CI: 0.49; 0.85) for dimethyl fumarate 240 mg two times daily, 0.68 (0.52; 0.89) for dimethyl fumarate 240 mg three times daily, 0.71 (0.55; 0.90) for fingolimod oral 0.5 mg, and 0.71 (0.56; 0.90) for fingolimod oral 1.25 mg.

Table 7. Relative risk for disability progression for active MS treatments compared to placebo

	Direct evide	nce	Indirect evi	lence	Network meta-	analysis
Interventions	RR (95% CI)	GRADE	RR (95% CI)	GRADE	RR (95% CI)	GRADE
Interferon beta-1a 22 mcg	0.84 (0.61 to 1.19)	Low	NA	NA	0.84 (0.61 to 1.19)	Low
Interferon beta-1a 30 mcg	0.68 (0.50 to 0.95)	Moderate	0.88 (0.66 to 1.20)	Low	0.8 (0.65 to 0.99)	Moderate
Interferon beta-1a 44 mcg	0.70 (0.48 to 1.04)	Low	0.86 (0.59 to 1.30)	Low	0.77 (0.6 to 1.01)	Low
Glatiramer acetate 20mg	0.88 (0.61 to 1.21)	Low	0.70 (0.51 to 0.94)	Low	0.78 (0.63 to 0.96)	Low
Dimethyl fumarate 240 mg two times daily	0.65 (0.49 to 0.85)	High	NA	NA	0.65 (0.49 to 0.85)	High
Dimethyl fumarate 240 mg three times daily	0.68 (0.52 to 0.89)	High	NA	NA	0.68 (0.52 to 0.89)	High
Teriflunomide oral 7 mg	0.8 (0.55 to 1.13)	Low	NA	NA	0.8 (0.55 to 1.13)	Low
Teriflunomide oral 14 mg	0.73 (0.51 to 1.05)	Low	NA	NA	0.73 (0.51 to 1.05)	Low
Fingolimod oral 0.5 mg	0.75 (0.56 to 0.98)	High	0.56 (0.32 to 0.91)	Low	0.71 (0.55 to 0.9)	High
Fingolimod oral 1.25 mg	0.70 (0.52 to 0.92)	High	0.81 (0.48 to 1.31)	Low	0.71 (0.56 to 0.9)	High
Peginterferon beta-1a 125 mcg once every 2 wks	0.61 (0.36 to 0.98)	Low	NA	NA	0.61 (0.36 to 0.98)	Low
Peginterferon beta-1a 125 mcg once every 4 wks	0.62 (0.38 to 1.01)	Low	NA	NA	0.62 (0.38 to 1.01)	Low
Natalizumab	0.59 (0.42 to 0.84)	Moderate	NA	NA	0.59 (0.42 to 0.84)	Moderate
Interferon beta-1b 250 mcg	0.77 (0.50 to 1.17)	Low	0.67 (0.43 to 0.95)	Low	0.72 (0.54 to 0.92)	Low
Alemtuzumab 12 mg IV q.d	NA	NA	0.4 (0.27 to 0.6)	Low	0.4 (0.27 to 0.6)	Low
Alemtuzumab 24 mg IV q.d	NA	NA	0.36 (0.16 to 0.74)	Very low	0.36 (0.16 to 0.74)	Very low
Interferon beta-1b 500 mcg SC 1/2 d.	NA	NA	0.79 (0.56 to 1.1)	Low	0.79 (0.56 to 1.1)	Low

RR= relative ratio, CI= confidence interval, mg= milligrams, mcg= micrograms, SC= subcutaneous, IM= intra muscular, q.d.= once daily, q.w.=once weekly, t.i.w.= three times weekly, 2.i.d= two times daily, t.i.d= three times daily, 1/2 w=once every 2 weeks, 1/4 w=once every 4 weeks.

Active treatments compared with each other

We obtained similar results when comparing active treatments with each other using direct and indirect evidence, and the evidence from the whole network (except for interferon beta-1b 250 mcg versus interferon beta-1a 30 mcg) (Table 8). We had evidence of very low to moderate quality (Table 8). Only two of the network meta-analysis comparisons showed statistically significant differences between treatments. interferon beta-1a 44 mcg was less effective against disability progression than alemtuzumab 12 mg and 24 mg, with RRs of 1.95 (95% CI: 1.45; 2.59) (evidence of moderate quality) and 2.15 (1.10; 4.55) (evidence of very low quality), respectively.

		Direct evi	dence	Indirect evid	lence	Network Meta-a	analysis
Intervention	Comparison	RR (95% CI)	GRADE	RR (95% CI)	GRADE	RR (95% CI)	GRADE
Alemtuzumab 24 mg	Alemtuzumab 12 mg	0.85 (0.4 to 1.65)	Very low	NA	NA	0.85 (0.4 to 1.65)	Very low
Interferon beta-1a 44 mcg	Alemtuzumab 12 mg	1.95 (1.45 to 2.59)	Moderate	NA	NA	1.95 (1.45 to 2.59)	Moderate
Interferon beta-1a 44 mcg	Alemtuzumab 24 mg	2.15 (1.1 to 4.55)	Very low	NA	NA	2.15 (1.1 to 4.55)	Very low
Interferon beta-1a 44 mcg	Interferon beta-1a 22 mcg	0.92 (0.65 to 1.3)	Low	NA	NA	0.92 (0.65 to 1.3)	Low
Interferon beta-1a 44 mcg	Interferon beta-1a 30 mcg	0.89 (0.55 to 1.38)	Low	1.04 (0.72 to 1.50)	Low	0.97 (0.73 to 1.3)	Low
Interferon beta-1a 60 mcg	Interferon beta-1a 30 mcg	0.99 (0.71 to 1.39)	Low	NA	NA	0.99 (0.71 to 1.39)	Low
glatiramer acetate 20mg	Interferon beta-1a 30 mcg	1.18 (0.81 to 1.75)	Low	0.87 (0.64 to 1.17)	Low	0.98 (0.76 to 1.23)	Low
Fingolimod oral 0.5 mg	Interferon beta-1a 30 mcg	0.72 (0.42 to 1.17)	Low	0.96 (0.68 to 1.33)	Low	0.89 (0.65 to 1.16)	Low
Fingolimod oral 1.25 mg	Interferon beta-1a 30 mcg	0.99 (0.58 to 1.60)	Low	0.85 (0.59 to 1.19)	Low	0.89 (0.66 to 1.18)	Low
Interferon beta-1b 250 mcg	Interferon beta-1a 30 mcg	0.44 (0.23 to 0.82)	Low	1.07 (0.81 to 1.43)	Low	0.9 (0.65 to 1.17)	Low
glatiramer acetate 20mg	Interferon beta-1a 44 mcg	0.75 (0.46 to 1.21)	Low	1.17 (0.82 to 1.65)	Low	1.01 (0.75 to 1.33)	Low
dimethyl fumarate 240 mg two times daily	glatiramer acetate 20mg	0.78 (0.52 to 1.18)	Low	0.80 (0.51 to 1.18)	Low	0.83 (0.61 to 1.15)	Low
dimethyl fumarate 240 mg three times daily	glatiramer acetate 20mg	0.79 (0.53 to 1.16)	Low	0.88 (0.59 to 1.36)	Low	0.88 (0.64 to 1.18)	Low
Interferon beta-1b 250 mcg	glatiramer acetate 20mg	1.04 (0.74 to 1.46)	Moderate	0.74 (0.48 to 1.09)	Low	0.92 (0.69 to 1.16)	Moderate
Interferon beta-1b 500 mcg	glatiramer acetate 20mg	1.01 (0.74 to 1.36)	Moderate	NA	NA	1.01 (0.74 to 1.36)	Moderate
dimethyl fumarate 240 mg three times daily	dimethyl fumarate 240 mg two times daily	1.06 (0.78 to 1.42)	Low	NA	NA	1.06 (0.78 to 1.42)	Low
Teriflunomide oral 14 mg	Teriflunomide oral 7 mg	0.92 (0.64 to 1.35)	Low	NA	NA	0.92 (0.64 to 1.35)	Low
Fingolimod oral 1.25 mg	Fingolimod oral 0.5 mg	1.01 (0.78 to 1.32)	Moderate	NA	NA	1.01 (0.78 to 1.32)	Moderate
Peginterferon beta-1a 125 mcg once every 4 wks	Peginterferon beta-1a 125 mcg once every 2 wks	1.02 (0.61 to 1.74)	Low	NA	NA	1.02 (0.61 to 1.74)	Low
Interferon beta-1b 500 mcg	Interferon beta-1b 250 mcg	1.1 (0.84 to 1.51)	Moderate	NA	NA	1.1 (0.84 to 1.51)	Moderate

Table 8. Relative risk for disability progression for active MS treatments compared to others for comparisons with available direct evidence

RR= relative ratio, CI= confidence interval, mg= milligrams, mcg= micrograms, SC= subcutaneous, IM= intra muscular, q.d.= once daily, q.w.=once weekly, t.i.w.= three times

weekly, 2.i.d= two times daily, t.i.d= three times daily, 1/2 w=once every 2 weeks, 1/4 w=once every 4 weeks.

Withdrawal due to adverse events

We present here the results obtained using the "network meta-analysis approach". Those are consistent with results found with the "pairwise comparison method". The "pairwise comparison method" results are presented in Appendix 7.

Figure 5 presents the network of evidence available for the outcome withdrawal due to adverse events. This network included information on 19 different treatments strategies and placebo.

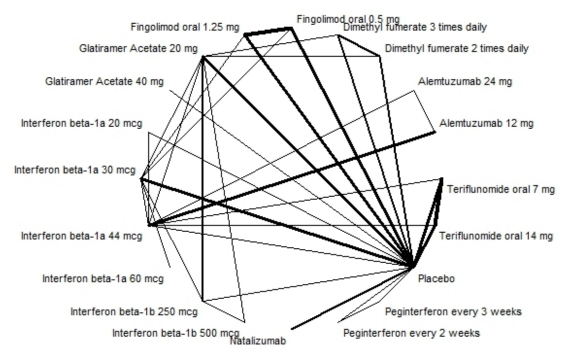


Figure 5. Evidence network for withdrawal due to adverse events

Active treatments versus placebo

Table 9 presents results estimated through direct and indirect evidence, and through the whole network. Results are consistent (except for interferon beta-1b 250 mcg). We had evidence for 19 treatments versus placebo. The quality of the evidence considered for the whole network was of very low to moderate quality. Four treatments were statistically significantly more associated with withdrawal due to adverse events than placebo. We found RRs for withdrawal due to adverse events of 2.20 (95% CI: 1.29-3.97) for interferon beta-1a 44 mcg (low quality evidence), of 2.21 (1.42; 3.58) for fingolimod oral 1.25 mg (moderate quality), and of 3.57 (1.27; 11.14) and 3.47 (1.25 to 10.9) for peg-interferon beta-1a 125 mcg once every 2 and 4 weeks, respectively (low quality evidence).

Table 9. Relative risk for withdrawal da	ue to adverse events for active MS trea	tments compared to placebo

	Direct evidence		Indiret evidence		Network meta-analysis	
Interventions	RR (95% CI)	GRADE	RR (95% CI)	GRADE	RR (95% CI)	GRADE
Interferon beta-1a 22 mcg	1.68 (0.5 to 5.98)	Low	NA	NA	1.68 (0.5 to 5.98)	Low
Interferon beta-1a 30 mcg	1.73 (0.82 to 3.87)	Low	1.12 (0.61 to 2.10)	Low	1.33 (0.85 to 2.17)	Low
Interferon beta-1a 44 mcg	5.32 (1.09 to 41.63)	Low	1.98 (1.10 to 3.61)	Low	2.2 (1.29 to 3.97)	Low
Glatiramer acetate 20mg	1.22 (0.64 to 2.66)	Low	1.15 (0.54 to 2.42)	Low	1.17 (0.74 to 1.94)	Low
Glatiramer acetate 40mg	2.5 (0.86 to 8.29)	Low	NA	NA	2.5 (0.86 to 8.29)	Low
Dimethyl fumarate 240 mg 2.i.d.	1.24 (0.74 to 2.13)	Low	NA	NA	1.24 (0.74 to 2.13)	Low
Dimethyl fumarate 240 mg t.i.d	1.25 (0.74 to 2.13)	Low	NA	NA	1.25 (0.74 to 2.13)	Low
Teriflunomide oral 7 mg	1.54 (0.89 to 2.51)	Low	0.89 (0.32 to 2.44)	Low	1.37 (0.82 to 2.21)	Low
Teriflunomide oral 14 mg	1.70 (1.02 to 3.01)	Low	1.29 (0.47 to 3.44)	Low	1.53 (0.96 to 2.54)	Low
Fingolimod oral 0.5 mg	1.49 (0.86 to 2.50)	Low	1.48 (0.65 to 3.55)	Low	1.54 (0.98 to 2.52)	Low
Fingolimod oral 1.25 mg	1.93 (1.18 to 3.14)	Moderate	3.26 (1.52 to 7.22)	Low	2.21 (1.42 to 3.58)	Moderate
Peginterferon beta-1a 125 mcg 1/2 w	3.57 (1.27 to 11.14)	Low	NA	NA	3.57 (1.27 to 11.14)	Low
Peginterferon beta-1a 125 mcg 1/4 w	3.47 (1.25 to 10.9)	Low	NA	NA	3.47 (1.25 to 10.9)	Low
Natalizumab	1.22 (0.5 to 2.74)	Low	NA	NA	1.22 (0.5 to 2.74)	Low
Interferon beta-1b 250 mcg	0.07 (0.003 to 0.48)	Low	1.64 (0.68 to 4.36)	Low	0.84 (0.4 to 1.87)	Low
Alemtuzumab 24 mg IV q.d	NA	NA	0.54 (0.17 to 1.54)	Very low	0.54 (0.17 to 1.54)	Very low
Alemtuzumab 12 mg IV q.d	NA	NA	0.61 (0.25 to 1.47)	Low	0.61 (0.25 to 1.47)	Low
Interferon beta-1b 500 mcg SC 1/2 d	NA	NA	1.37 (0.52 to 3.92)	Low	1.37 (0.52 to 3.92)	Low
Interferon beta-1a 60 mcg IM q.w	NA	NA	1.9 (0.79 to 4.81)	Low	1.9 (0.79 to 4.81)	Low

RR= relative ratio, CI= confidence interval, mg= milligrams, mcg= micrograms, SC= subcutaneous, IM= intra muscular, q.d.= once daily, q.w.=once weekly, t.i.w.= three times weekly, 2.i.d= two times daily, t.i.d= three times daily, 1/2 w=once every 2 weeks, 1/4 w=once every 4 weeks.

Active treatments compared with each other

Results using direct and indirect evidence, and evidence from the whole network were similar in terms of direction of the association and magnitude (Table 10). The quality of the evidence ranged from very low to moderate. Only two of the network meta-analysis comparisons showed statistically significant results. Patients withdrew more due to adverse events with interferon beta-1a 44 mcg than with alemtuzumab 12 and 24 mg (RRs of 3.6 (95% CI: 1.88; 7.33), and 4.08 (1.69; 11.42), respectively). The corresponding quality of the evidence was moderate and very low.

Table 10. Relative risk for withdrawal due to adverse events for active MS treatments compo	ared to each other

		Directe evidence		Indirecte evidence		Network meta-analysis	
Intervention	Comparison	RR (95% CI)	GRADE	RR (95% CI)	GRADE	RR (95% CI)	GRADE
Alemtuzumab 24 mg	Alemtuzumab 12 mg	0.88 (0.3 to 2.31)	Low	NA	NA	0.88 (0.3 to 2.31)	Low
Interferon beta-1a 44 mcg	Alemtuzumab 12 mg	3.6 (1.88 to 7.33)	Moderate	NA	NA	3.6 (1.88 to 7.33)	Moderate
Interferon beta-1a 44 mcg	Alemtuzumab 24 mg	4.08 (1.69 to 11.42)	Very low	NA	NA	4.08 (1.69 to 11.42)	Very low
Interferon beta-1a 44 mcg	Interferon beta-1a 22 mcg	1.31 (0.4 to 4.36)	Low	NA	NA	1.31 (0.4 to 4.36)	Low
Interferon beta-1a 44 mcg	Interferon beta-1a 30 mcg	1.15 (0.43 to 3.10)	Low	2.09 (0.98 to 4.57)	Low	1.65 (0.91 to 3.08)	Low
Interferon beta-1a 60 mcg	Interferon beta-1a 30 mcg	1.43 (0.66 to 3.11)	Low	NA	NA	1.43 (0.66 to 3.11)	Low
glatiramer acetate 20mg	Interferon beta-1a 30 mcg	0.61 (0.22 to 1.67)	Low	1.02 (0.53 to 2.03)	Low	0.88 (0.51 to 1.55)	Low
Fingolimod oral 0.5 mg	Interferon beta-1a 30 mcg	1.28 (0.52 to 3.44)	Low	1.17 (0.58 to 2.29)	Low	1.16 (0.65 to 2.04)	Low
Fingolimod oral 1.25 mg	Interferon beta-1a 30 mcg	2.44 (1.09 to 5.68)	Low	1.41 (0.73 to 2.59)	Low	1.66 (0.94 to 2.91)	Low
Interferon beta-1b 250 mcg	Interferon beta-1a 30 mcg	6.27 (0.79 to 172.3)	Low	0.41 (0.16 to 0.93)	Low	0.63 (0.28 to 1.44)	Low
glatiramer acetate 20mg	Interferon beta-1a 44 mcg	0.88 (0.36 to 1.94)	Low	0.37 (0.17 to 0.77)	Low	0.53 (0.29 to 0.96)	Low
Teriflunomide oral 7 mg	Interferon beta-1a 44 mcg	0.40 (0.14 to 1.00)	Low	0.75 (0.34 to 1.42)	Low	0.62 (0.31 to 1.12)	Low
Teriflunomide oral 14 mg	Interferon beta-1a 44 mcg	0.54 (0.20 to 1.38)	Low	0.76 (0.35 to 1.57)	Low	0.69 (0.37 to 1.28)	Low
dimethyl fumarate 240 mg two times daily	glatiramer acetate 20mg	1.18 (0.49 to 2.84)	Low	0.96 (0.37 to 2.36)	Low	1.07 (0.56 to 1.92)	Low
dimethyl fumarate 240 mg three times daily	glatiramer acetate 20mg	1.15 (0.52 to 2.56)	Low	0.98 (0.35 to 2.53)	Low	1.07 (0.56 to 1.93)	Low
Interferon beta-1b 250 mcg	glatiramer acetate 20mg	0.91 (0.37 to 2.27)	Low	0.49 (0.14 to 1.63)	Low	0.72 (0.35 to 1.49)	Low
Interferon beta-1b 500 mcg	glatiramer acetate 20mg	1.16 (0.46 to 3.05)	Low	NA	NA	1.16 (0.46 to 3.05)	Low
dimethyl fumarate 240 mg three times daily	dimethyl fumarate 240 mg two times daily	1.01 (0.58 to 1.73)	Low	NA	NA	1.01 (0.58 to 1.73)	Low
Teriflunomide oral 14 mg	Teriflunomide oral 7 mg	1.12 (0.73 to 1.85)	Moderate	NA	NA	1.12 (0.73 to 1.85)	Moderate
Fingolimod oral 1.25 mg	Fingolimod oral 0.5 mg	1.43 (0.94 to 2.21)	Moderate	NA	NA	1.43 (0.94 to 2.21)	Moderate
Peginterferon beta-1a 125 mcg once every 4 wks	Peginterferon beta-1a 125 mcg once every 2 wks	0.98 (0.41 to 2.37)	Low	NA	NA	0.98 (0.41 to 2.37)	Low
Interferon beta-1b 500 mcg	Interferon beta-1b 250 mcg	1.63 (0.66 to 4.11)	Low	NA	NA	1.63 (0.66 to 4.11)	Low

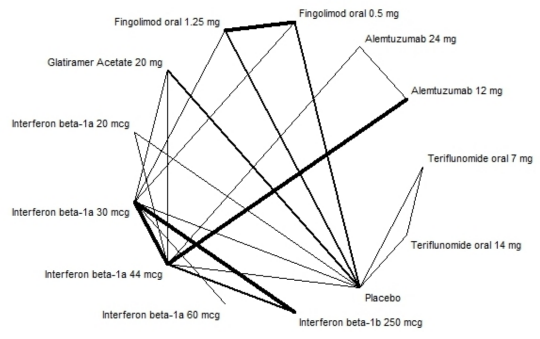
RR= relative ratio, CI= confidence interval, mg= milligrams, mcg= micrograms, SC= subcutaneous, IM= intra muscular, q.d.= once daily, q.w.=once weekly, t.i.w.= three times

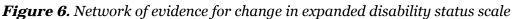
weekly, 2.i.d= two times daily, t.i.d= three times daily, 1/2 w=once every 2 weeks, 1/4 w=once every 4 weeks.

Change in Expanded Disability Scale

Here, we examined disability progression in a continuous manner; that is by estimating the change in EDSS. We did not grade the quality of the evidence for this outcome. We present here results for active treatments versus placebo. We compare results obtained though the "network meta-analysis approach" and the "pairwise comparison method".

The network of the evidence for change in EDSS included 12 treatment strategies and placebo (Figure 6).





Active treatments versus placebo

Twelve different treatments were compared to placebo in the network meta-analysis (Table 11). Four treatments were statistically significantly more effective than placebo against disability progression: alemtuzumab 24 mg (mean difference=-0.91 (95% CI:- 1.48; -0.4), alemtuzumab 12 mg (-06 (-1.02; -0.24)), interferon beta-1b 250 mcg every other day (-0.58 (-0.94; -0.22)), and interferon beta-1a 44 mcg three times a week (- 0.28 (-0.58; -0.02).

When comparing results obtained through "network meta-analysis approach" and "pairwise comparison method", we found a difference in the magnitude and statistical significance of the effect for the comparison interferon beta-1a 30 mcg versus placebo (Table 11). The mean difference in change in EDSS score was -0.59 (-0.86 to -0.32) when considering pairwise comparisons, and -0.22 (-0.48 to 0.02) for the network meta-analysis estimates. For the other treatments strategies, a similar magnitude of effect was seen.

Table 11. Change in expanded disability status scale for MS treatments compared to placebo for direct pairwise comparisons and network evidence

	Network meta-analysis		Pairwise comparison
Interventions	Mean difference	SUCRA	Mean difference
Alemtuzumab 24 mg IV q.d	-0.91 (-1.48 to -0.4)	0.98	
Alemtuzumab 12 mg IV q.d	-0.6 (-1.02 to -0.24)	0.86	
Interferon beta-1b 250 mcg SC every other day	-0.58 (-0.94 to -0.22)	0.85	
Interferon beta-1a 44 mcg SC t.i.w	-0.28 (-0.58 to -0.02)	0.56	-0.24 (-0.48 to 0.00)
Interferon beta-1a 22 mcg SC t.i.w	-0.27 (-0.71 to 0.15)	0.52	-0.25 (-0.51 to 0.01)
Interferon beta-1a 60 mcg IM q.w	-0.25 (-0.76 to 0.24)	0.49	
Fingolimod oral 1.25 mg	-0.22 (-0.47 to 0.04)	0.46	-0.15 (-0.25 to -0.05)
Interferon beta-1a 30 mcg IM q.w	-0.22 (-0.48 to 0.02)	0.46	-0.59 (-0.86 to -0.32)
Teriflunomide oral 14 mg	-0.14 (-0.56 to 0.27)	0.35	-0.14 (-0.27 to -0.01)
Fingolimod oral 0.5 mg	-0.16 (-0.41 to 0.1)	0.35	-0.08 (-0.20 to 0.03)
glatiramer acetate 20mg q.d	-0.13 (-0.4 to 0.11)	0.31	-0.03 (-0.12 to 0.06)
Teriflunomide oral 7 mg	-0.05 (-0.47 to 0.36)	0.23	-0.05 (-0.18 to 0.08)
Placebo	0	0.10	

CI= confidence interval, mg= milligrams, mcg= micrograms, SC= subcutaneous, IM= intra muscular, q.d.= once daily, q.w.=once weekly, t.i.w.= three times weekly, 2.i.d= two times daily, t.i.d= three times daily, 1/2 w=once every 2 weeks, 1/4 w=once every 4 weeks. SUCRA= surface under the cumulative ranking curve.

Serious adverse events

We present here results for active treatments versus placebo. We did not grade the quality of the evidence for this outcome. We compare results obtained though the "network meta-analysis approach" and the "pairwise comparison method".

The evidence network available for serious adverse events is presented in Figure 7.

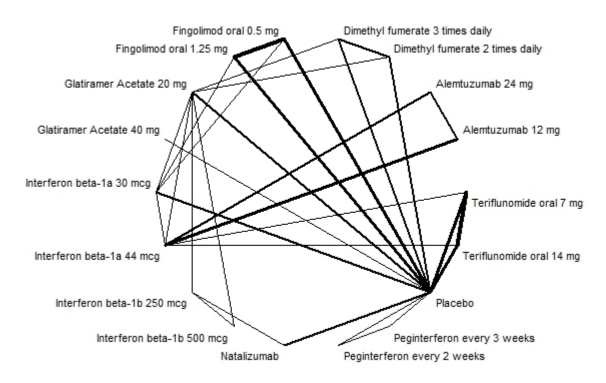


Figure 7. Network of evidence for serious adverse events

Active treatments versus placebo

Through the network meta-analysis, we had information for 17 treatments (Table 12). When considering all the available evidence comparing active treatments and placebo, based on the confidence intervals, no statistically significant difference was seen between results obtained through pairwise comparisons and network meta-analysis results. However, for the network meta-analysis results no treatments were found to increase statistically significantly serious adverse events compared to placebo. Results from the "pairwise comparison method" showed that peg-interferon beta-1a 125 mcg once every 4 and 2 weeks were associated with more serious adverse events than placebo, with RRs of 1.55 (95% CI: 1.12-2.14) and 1.66 (1.21- 2.28), respectively.

Table 12. Relative risk for serious adverse events for MS treatments compared to placebo for direct pairwise comparisons and network evidence

	Network meta-analysis		Pairwise comparison
Intervention	Relative ratio (95% CI)	SUCRA	Relative ratio (95% CI)
Alemtuzumab 12 mg IV q.d	0.67 (0.37 to 1.28)	0.80	
Interferon beta-1b 250 mcg SC every other day	0.66 (0.35 to 1.26)	0.80	
Dimethyl fumarate 240 mg three times daily	0.72 (0.49 to 1.07)	0.76	0.73 (0.59 to 0.91)
Interferon beta-1a 30 mcg IM q.w	0.77 (0.54 to 1.13)	0.70	0.65 (0.44 to 0.97)
Glatiramer acetate 20mg q.d	0.78 (0.54 to 1.14)	0.68	0.99 (0.50 to 1.97)
Alemtuzumab 24 mg IV q.d	0.79 (0.42 to 1.53)	0.64	
Dimethyl fumarate 240 mg two times daily	0.81 (0.56 to 1.19)	0.64	0.82 (0.67 to 1.01)
Natalizumab	0.81 (0.49 to 1.39)	0.62	0.80 (0.62 to 1.03)
Interferon beta-1a 44 mcg SC t.i.w	0.86 (0.52 to 1.46)	0.54	
Interferon beta-1b 500 mcg SC every other day	0.93 (0.49 to 1.8)	0.47	
Fingolimod oral 0.5 mg	0.96 (0.68 to 1.39)	0.45	0.98 (0.67 to 1.42)
Glatiramer acetate 40mg t.i.w	0.99 (0.49 to 2.04)	0.44	0.98 (0.59 to 1.63)
Placebo	1	0.39	
Teriflunomide oral 7 mg	1.03 (0.71 to 1.51)	0.37	1.02 (0.79 to 1.32)
Teriflunomide oral 14 mg	1.07 (0.73 to 1.54)	0.33	1.14 (0.89 to 1.46)
Fingolimod oral 1.25 mg	1.22 (0.87 to 1.77)	0.20	1.18 (0.73 to 1.91)
peginterferon beta-1a 125 mcg once every 4 weeks	1.55 (0.88 to 2.74)	0.11	1.55 (1.12 to 2.14)
Peginterferon beta-1a 125 mcg once every 2 weeks	1.67 (0.94 to 2.94)	0.07	1.66 (1.21 to 2.28)

RR= relative ratio, CI= confidence interval, mg= milligrams, mcg= micrograms, SC= subcutaneous, IM= intra muscular, q.d.= once daily, q.w.=once weekly, t.i.w.= three times weekly, 2.i.d= two times daily, t.i.d= three times daily, 1/2 w=once every 2 weeks, 1/4 w=once every 4 weeks. SUCRA= surface under the cumulative ranking curve.

Mortality

We present here results for active treatments versus placebo. We compare results obtained though the "network meta-analysis approach" and the "pairwise comparison method". Figure 8 illustrates the network of evidence available for mortality. In total, 19 treatment strategies and placebo were examined.

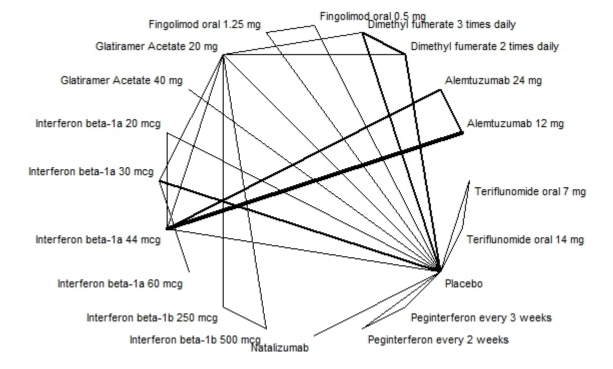


Figure 8. Evidence network for mortality

Active treatments versus placebo

Table 13 reports results for nineteen treatments compared to placebo. Estimates obtained through "pairwise comparison method" and "network meta-analysis approach" are statistically consistent. None of the examined treatments were associated with a higher risk for mortality than placebo. **Table 13.** Relative risk for mortality for MS treatments compared to placebo for direct pairwise comparisons and network evidence

	Network meta-analysis		Pairwise comparison
Intervention	Relative ratio (95% CI)	SUCRA	Relative ratio (95% CI)
Fingolimod oral 0.5 mg	0.1 (0. to 2.57)	0.80	0.20 (0.01 to 4.09)
Interferon beta-1b 500 mcg SC every other day	0.08 (0. to 5.9)	0.79	
Glatiramer acetate 40mg t.i.w	0.08 (0. to 3.54)	0.79	0.16 (0.01 to 4.00)
Interferon beta-1b 250 mcg SC every other day	0.07 (0. to 6.65)	0.79	
Peginterferon beta-1a 125 mcg once every 4 weeks	0.4 (0.01 to 10.22)	0.61	0.50 (0.05 to 5.50)
Peginterferon beta-1a 125 mcg once every 2 weeks	0.41 (0.01 to 8.87)	0.61	0.49 (0.04 to 5.37)
Dimethyl fumarate 240 mg two times daily	0.52 (0.04 to 5.34)	0.59	1.00 (0.10 to 9.62)
Fingolimod oral 1.25 mg	0.52 (0.02 to 6.76)	0.58	0.49 (0.04 to 5.35)
Dimethyl fumarate 240 mg three times daily	0.89 (0.09 to 8.41)	0.47	1.64 (0.20 to 13.27)
Interferon beta-1a 44 mcg SC t.i.w	0.97 (0.06 to 17.15)	0.47	0.34 (0.01 to 8.26)
Glatiramer acetate 20mg q.d	0.9 (0.11 to 7.85)	0.47	1.03 (0.06 to 16.47)
Teriflunomide oral 14 mg	0.94 (0.02 to 37.74)	0.46	0.94 (0.06 to 15.00)
Placebo	1	0.44	
Interferon beta-1a 22 mcg SC t.i.w	1.6 (0.07 to 34.77)	0.37	0.99 (0.06 to 15.70)
Alemtuzumab 24 mg IV q.d	2.08 (0.04 to 125.5)	0.34	
Interferon beta-1a 60 mcg IM q.w	2.28 (0.03 to 222.1)	0.34	
Interferon beta-1a 30 mcg IM q.w	2.1 (0.26 to 24.45)	0.29	2.86 (0.30 to 27.43)
Teriflunomide oral 7 mg	2.59 (0.12 to 82.51)	0.29	2.08 (0.19 to 22.79)
Alemtuzumab 12 mg IV q.d	2.81 (0.08 to 168.2)	0.27	
Natalizumab	4.34 (0.16 to 2761.)	0.22	2.52 (0.12 to 52.25)

RR= relative ratio, CI= confidence interval, mg= milligrams, mcg= micrograms, SC= subcutaneous, IM= intra muscular, q.d.= once daily, q.w.=once weekly, t.i.w.= three times weekly, 2.i.d= two times daily, t.i.d= three times daily, 1/2 w=once every 2 weeks, 1/4 w=once every 4 weeks. SUCRA= surface under the cumulative ranking curve.

Stay at hospitals

Very few studies reported on stay at hospitals. Therefore, we could not summarise quantitatively the results for this endpoint.

Economic evaluation-Methods

General

In order to assess the health economic effectiveness of different disease-modifying medicines for patients with RRMS, we performed a cost-utility analysis (CUA). The relevant costs were expressed in 2015 Norwegian kroner (NOK), and effects were expressed in quality-adjusted life-years (QALYs). Both costs and effects were discounted using an annual discount rate of 4% as recommended by the Norwegian Ministry of Finance and guidelines for health economic evaluation in the health sector (67).

The analysis was carried out from a healthcare perspective. The healthcare perspective is relevant for prioritisation of interventions within a fixed budget if the aim of the decision maker is to maximize health (no expansion of the budget is assumed). The methodological guidelines for economic evaluation in the health sector recommend a societal perspective that includes consequences for all parts of the economy, including time costs, the deadweight loss of taxation, any productivity changes, and excluding transfers such as value added tax. This perspective is more appropriate if an expansion of the budget is assumed and in settings where prioritization of interventions across sectors of the economy is relevant (e.g. for public health interventions).

We expressed the results as mean incremental cost-effectiveness ratio (ICER) from 10,000 runs of the model in base-case. We handled uncertainties in model parameters by performing probabilistic sensitivity analyses, designed as a Monte Carlo simulation, with 10,000 iterations.

Population, interventions and model structure

Population

In the economic evaluation, we assumed that a typical RRMS patient population in Norway has an average age of 30 years at diagnosis, and 68% are female.

Interventions

There are currently 12 disease-modifying therapies approved and available for RRMS patients in Norway (based on clinical experts' opinion). All these active treatment options were included in our analysis (Table 14).

Interventions
Alemtuzumab 12 mg (Lemtrada)
Dimethyl fumarate 240 mg (Tecifidera)
Fingolimod 0.5 mg (Gilenya)
Glatiramer acetate 20 mg (Copaxone)*
Interferon beta-1a 30 mcg (Avonex)
Interferon beta-1a 22 mcg (Rebif)
Interferon beta-1a 44 mcg (Rebif)
Interferon beta-1b 250 mcg (Betaferon)
Interferon beta-1b 250 mcg (Extavia)
Natalizumab 300 mg/15 mL (Tysabri)
Peg-interferon beta-1a 125 mcg (Plegridy)
Teriflunomide 14 mg (Aubagio)

mg: milligram; mL: millilitre; mcg: microgram

* Glatiramer acetate 40 mg 3 times per week was discussed in the discussion section.

Because of lack of clinical data exploring the sequential use of different treatment options following the failure of first-line treatments or switching, we assumed that patients could not switch between treatments in the model.

Model structure

In order to assess the cost-utility of different disease-modifying therapies in patients diagnosed with RRMS, a decision analytic model was developed in TreeAge pro ® 2015. The model is of the Markov type, in which a cohort of patients is followed over a given period of time. A Markov model was considered appropriate, as multiple sclerosis is a chronic condition requiring continuous treatment (68, 69).

We developed the model based on a previously published report with similar objectives as ours (27). The validity of the model structure and assumptions to the Norwegian context have been discussed and evaluated by two independent clinical experts experienced in treating patients with RRMS in Norway. The model structure and all assumptions were adapted to the Norwegian setting, and took into consideration Norwegian clinical practice.

The model simulates the natural history of MS using the state transition methodology (Figure 9). Health states were defined according to the Kurtzke EDSS (70). EDSS is a clinical rating scale ranging from 0 to 10. EDSS 0-2.5 refers to patients with no or few limitations in mobility, and EDSS 10 refers to death due to MS. Disability status was modelled from 0 to 10 for RRMS and from 2 to 10 for SPMS (70).

During one cycle, all patients could remain in the current health state, progress to the next more severe state, transition to a secondary-progressive health state, or die (Figure 9). Patients with an EDDS scale of five or lower could also improve to a less severe state, and stop treatment. Improvement in lower health states was modelled by assuming that a maximum of 2 EDSS-point improvements could be achieved (71). Patients would discontinue treatment once they progress to an EDSS of six or SPMS (based on clinical experts' opinion).

In the base-case analysis, we assumed no treatment effect once patients progress to an EDDS of six. It is also documented that with advancing disease (EDSS>6) less relapses occur (71). We, therefore, assumed that relapses would occur only in patients with EDSS of five or lower.

We assessed the costs and utilities associated with different treatment options over 20 years for the base case analysis (based on experts' opinion). Alternatives horizons of 10 years and 30 years were considered in scenario analyses. We used a cycle length of the model of one year, meaning that any transitions between different states could happen only once a year. Patients could be in only one of the pre-defined states at any time. Upon completion of each cycle, patients could, depending on transition probabilities, transfer to another state or remain in the same state until death or the end of the simulation. Each state and event is associated with specific health outcomes and costs. Death is modelled as an absorbing state. Once an individual makes a transition into the absorbing state, no further incurred costs or health outcome are included in the analysis.

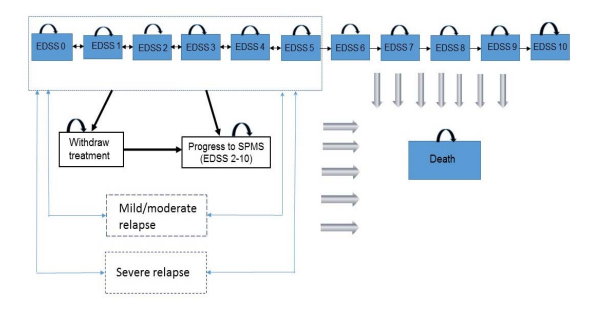


Figure 9. Model structure

EDSS: Expanded Disability Status Scale; RRMS: relapsing-remitting multiple sclerosis; SMPM: Secondary-progressive multiple sclerosis Note: Patients with EDSS over 5 can also progress to SPMS. Mild or moderate and severe relapses can occur in EDSS below 6 as events.

Disease-modifying therapies are usually initiated in patients with an EDSS score lower than 5, and mostly for patients with an EDSS score between 1 and 3 (clinical expert opinion and (72)). EDSS distributions used in our analysis are presented in Table 15.

EDSS score	Distributions (%)	Standard error
0	5.10	0.003
1	24.60	0.013
2	29.30	0.015
3	24.70	0.013
4	12.70	0.006
≥ 5	3.60	0.002

Table 15. EDSS distribution

EDSS: Expanded Disability Status Scale Source: Nixon *et.a*l 2014 (72)

Model Parameters

The model was created as a probabilistic model. This means that all uncertain parameters (efficacy, costs, epidemiological data, etc.) were modelled as probability distributions rather than point estimates. This was done to facilitate probabilistic sensitivity analysis. The sources and methods used to derive the model parameters are described below. First, we describe how we estimated the natural history transitional probabilities, then we describe how we incorporated into the model the clinical effect estimates (obtained through the systematic review (SR) and the network meta-analysis). Finally, we describe the methods used to calculate costs, and quality of life estimates.

Key model assumptions

Based on reporting of withdrawals in studies included in our SR, we set annual treatment discontinuation rate at 15% for the first two years in the base case analysis. This rate is also applicable to the Norwegian context according to the experts' opinion. A previous study showed that the proportion of patients who discontinued treatment and the degree of treatment adherence were similar across different treatment options (73). We therefore assumed the same discontinuation rate across all treatment options. We assumed no discontinuation after two years (expert opinion). Any patients who discontinues therapy subsequently progress according to natural history rates with no additional cost of therapy.

We assumed that treatments have no survival benefit. The annual risk of other mortality causes is, therefore, assumed to be the same as the normal population. We collected age and gender specific Norwegian all-cause mortality data from Statistics Norway (74). A weighted average was calculated based on the assumption that 68% of RRMS patients were female.

Natural history transitional probabilities

We did not find Norwegian data that were compatible to the developed model, so the transitional probabilities had to be based on estimates reported in the published literature. However, the transferability of the data to the Norwegian context were critically discussed and modified based on expert advice.

Disability progression

Probabilities for disability progression within RRMS health states, transitioning from RRMS to SPMS, as well as disability progression within SPMS health states were derived from a large 25- year patient-level cohort study (untreated patients) undertaken in London, Ontario, Canada (75, 76). The reported data were eligible for our model and used by the several previously published economic studies (27, 77).

Instantaneous hazard rates for disability progression without disease-modifying therapy were calculated from the Ontario dataset using the formula below (76), and are presented in the Tables 16-18.

 $\lambda \mathbf{i} = \frac{\text{Number of people leaving state i}}{\sum_{j=1}^{n} duration \text{ in state } \mathbf{i}}$

where n is the number of individuals, j is each individual leaving state i, and i= EDSS sate 0 to 10.

All rates were transformed into transition probabilities for use in the model (78). All natural history probabilities were incorporated in the model as beta distributions

EDSS score	Estimates (per person-year)	Variance
0	0.144	0.00007
1	0.075	0.00003
2	0.152	0.00006
3	0.272	0.00025
4	0.450	0.00166
5	0.485	0.00213
6	0.283	0.00104
7	0.342	0.00450
8	0.105	0.00139
9	0.167	0.02778

Table 16. Progression rates within RRMS health states

EDSS: Expanded Disability Status Scale Source: (27, 76)

Table 17. Progression rates from RRMS to SPMS

EDSS score	Estimates (per person-year)	Variance
0	0.004	0.000002
1	0.002	0.000001
2	0.029	0.000012
3	0.102	0.000094
4	0.199	0.000735
5	0.256	0.001126
6	0184	0.000676
7	0.237	0.000312
8	0.066	0.000866
9	0.167	0.027778

EDSS: Expanded Disability Status Scale; SPMS: Secondary-progressive multiple sclerosis Source: (27, 76)

Table 18. Progression rates within SPMS health states

EDSS score	Estimates (per person-year)	Variance
2	0.370	0.00370
3	0.385	0.00129
4	0.594	0.00280
5	0.349	0.00088
6	0.241	0.00029
7	0.186	0.00024
8	0.107	0.00015
9	0.093	0.00038

EDSS: Expanded Disability Status Scale; SPMS: Secondary-progressive multiple sclerosis Source: (27, 76)

Improvements in MS disability

Based on a large study, Tremlett and co-workers concluded that improvements in MS disability over one or two years were not unusual (71). The result of the study indicated that 8.3% of patients had an improvement of at least 1 point in the EDDS scale after one year, and 2.2% showed greater than or equal to 2-point improvements. We considered a maximum of two EDSS-point improvements in the model. The rates of annual disability improvements were used in the model only for the EDSS states lower than 6.

Relapse rate

There were no available Norwegian data on annual relapse rate compatible to our model. We considered therefore the best available sources. Annual relapse rates have been estimated based on Ontario cohort data (76), and published evidence suggested that the frequency of relapse is affected by a patient's age and disease duration (a decrease over time) (79, 80). Based on Ontario cohort data, the mean relapse rate after two years since disease onset was reported to be 0.835 and 1.423 for patients in EDSS 0 to 2 and 3+, respectively (76). These estimates were adjusted such that the patients enter the model with an average time since disease onset of five years and onwards (based on the studies included in our systematic review). More detailed information about the estimation of annual relapse rate can be found in the Canadian HTA report (27). These annual relapse rates were judged applicable to the Norwegian context by our clinical experts.

We used a Gamma distribution for annual relapse rates based on the assumption that events with a known average rate occur in a fixed interval of time.

Table 19. Annual relapse rates

Year since MS onset	Base estimate	Standard error					
For patients with a EDSS 0 to 2.5							
5	0.712	0.343					
10	0.623	0.335					
15	0.571	0.331					
20	0.534	0.327					
25	0.506	0.325					
For patients with	a EDSS 3 to 5.5						
5	1.255	0.386					
10	1.101	0.374					
15	1.011	0.367					
20	0.947	0.362					
25	0.897	0.358					

EDSS: Expanded Disability Status Scale Source: (27)

Based on published literature and expert opinion, we assumed that 23% of relapses were severe (81). In addition, we assumed that the average length of mild or moderate relapses was of 45 days. For severe relapse, it was of 90 days (27, 81).

Clinical efficacy parameters in the model

Clinical efficacy data for the model were the data presented in the "Clinical evaluationresults" section of this report. These were the results obtained through the network meta-analysis of the included trials. In the health economic model, we included the estimates on relapse rates and disability progression. These efficacy estimates were modelled by applying the relative risk for each treatment compared to best supportive care "no treatment", to the transitional probabilities based on the natural history of the disease for untreated patients.

We added the relative risks to the model as probability distributions. We used lognormal distributions, according to the methodology described by Briggs and co-authors (78). Standard errors for the log-normal distributions were calculated based on confidence intervals for efficacy estimates. The estimates of the calculations of distributions for efficacy parameters used in the model are presented in Tables 20 and 21. Based on expert opinion, we considered a reduction in treatment effect over time. Full effect of treatments is assumed to be 100% for the first four years, 75% from year 5 - 10, and 50% beyond 10 years.

Treatment effect on disability progression

The relative risks of sustained disability progression were multiplied to the transitional probabilities of patients moving to higher health states, as well as to progression to SPMS health states.

We assumed that patients transitioned as natural history of disease transitional probabilities between SPMS health state. That is treatments had no effect on the transition between SPMS states. Patients who withdraw treatment will progress according to transitional probabilities for natural disability progression, but will retain any previously accrued benefits.

Interventions	RR of sustained disability progression	Ln (RR)	SE
Alemtuzumab 12 mg (Lemtrada)	0.36	-1.02	0.39
Dimethyl fumarate 240 mg (Tecifidera)	0.65	-0.43	0.14
Fingolimod 0.5 mg (Gilenya)	0.71	-0.34	0.13
Glatiramer acetate 20 mg * (Copaxone)	0.78	-0.25	0.11
Interferon beta-1a 30 mcg (Avonex)	0.80	-0.22	0.11
Interferon beta-1a 22 mcg (Rebif)	0.84	-0.17	0.17
Interferon beta-1b 250 mcg (Betaferon)	0.72	-0.33	0.14
Interferon beta-1b 250 mcg (Extavia)	0.72	-0.33	0.14
Natalizumab 300 mg/15 mL (Tysabri)	0.59	-0.53	0.18
Peg-interferon beta-1a 125 mcg (Plegridy)	0.61	-0.49	0.26
Teriflunomide 14 mg (Aubagio)	0.73	-0.31	0.18

Table 20. Efficacy estimates for disability progression (log-normal distribution)

RR: relative risk; SE: standard error; mg: milligram; mL: millilitre; mcg: microgram

* We did not find any documentation for glatiramer acetate 40 mg.

Treatment effect on relapses

The expected number of relapses for each treatment alternative were estimated in the model by multiplying the treatment effect on the relapse rates for each treatment alternative (Table 21) to the average number of relapses experienced with "no treatment".

Interventions	RR of annual relapse rate	Ln (RR)	SE
Alemtuzumab 12 mg (Lemtrada)	0.29	-1.24	0.11
Dimethyl fumarate 240 mg (Tecifidera)	0.50	-0.69	0.09
Fingolimod 0.5 mg (Gilenya)	0.46	-0.78	0.08
Glatiramer acetate 20 mg * (Copaxone)	0.65	-0.43	0.05
Interferon beta-1a 30 mcg (Avonex)	0.82	-0.20	0.06
Interferon beta-1a 22 mcg (Rebif)	0.69	-0.37	0.10
Interferon beta-1a 44 mcg (Rebif)	0.64	-0.45	0.06
Interferon beta-1b 250 mcg (Betaferon)	0.66	-0.42	0.07
Interferon beta-1b 250 mcg (Extavia)	0.66	-0.42	0.07
Natalizumab 300 mg/15 mL (Tysabri)	0.30	-1.20	0.10
Peg-interferon beta-1a 125 mcg (Plegridy)	0.65	-0.43	0.14
Teriflunomide 14 mg (Aubagio)	0.67	-0.40	0.07

Table 21. Efficacy estimates for annual relapse (log-normal distribution)

RR: relative risk; SE: standard error; mg: milligram; mL: millilitre; mcg: microgram

* Glatiramer acetate 40 mg RR: 0.66 SE: 0.11

Treatment-related adverse events

Generally, disease-modifying therapies are well tolerated. Our systematic review showed no statistically significant differences between the therapies for serious adverse events. Moreover, most of the adverse events related to the RRMS treatments were transient, and some of them may potentially be related to the disease process (e.g. depression). We have therefore not included adverse events (except for Progressive multifocal leukoencephalopathy (PML)) in the model based on the assumption that the costs and disutility associated with adverse events would not have a significant impact on the results. However, some of the differences for resource use related to the adverse events have been considered when estimating of monitoring costs associated with each of the treatment strategies. For more information, see Appendix 8.

Natalizumab has been reported to be associated with the development of PML, which is a rare but serious infectious or inflammatory disease. PML is a viral infection (JC-virus) leading to inflammation and finally demyelination, often resulting in severe disability or death (82). A study from 2013 found a risk of developing PML of 2.84 cases per 1000 patients who received natalizumab for MS (83). It was also reported that 22% of the reported natalizumab-associated PML patients died (83). The costs and reduction in quality of life associated with PML is addressed in the next sections.

It should be mentioned that recently PML has also been reported in a small number of patients treated with other disease-modifying therapies, such as dimethyl fumurate and fingolimod. Due to insufficient data, we included PML only for natalizumab in the model.

Costs

An annual cost per patient associated with different treatment alternatives was calculated for each health state and event in the model. The costs included in the model are drug costs, monitoring costs associated with the use of drugs, costs related to MS patients care (excluding drugs) at different EDSS levels, and costs related to the treatments of relapses and PML.

All costs were measured in 2015 Norwegian kroner (NOK) (based on the consumer price index for the first four months of 2015 (74)). The uncertainty surrounding cost parameters were assessed by using gamma distribution.

Annual drug costs

Drug costs were calculated based on the maximum pharmacy retail prices that we received from the Drug procurement cooperation (LIS). The annual drug cost was estimated based on recommended doses (LIS), and are presented in Table 22.

Table 22. Drug costs per patient inclusive VAT

Drug	Dosage and recommended treatment regimen ^a	Dosage form ^a	LIS price (NOK) ª	Pills/ syringes per package ª	Annual drug cost (NOK)
Alemtuzumab (Lemtrada)	12 mg/1.2 ml per day for 5 days, 12 mg/1.2 ml per day for 3 days after one year (IV)	Vial	63,757.09	1	318,785 (5 days first year), 191,271 (3 days second year) ^b
Dimethyl fumarate (Tecifidera)	120 mgx2 for 7 days, 240mg x2 /dag	Capsule	3,256.12 (start package) 12,936.70	14 56	168,670
Fingolimod (Gilenya)	0.5 mg/day	Capsule	15,125.39	28	197,170
Glatiramer acetate (Copaxone) °	20mg/mL I syringe/day (SC)	Pre-filled Syringe	6,702.38	28	87,370
Interferon beta-1a (Avonex)	30 mcg/0.5 ml Once per week (IM)	Pre-filled Syringe	8,021.97	4	104,286
Interferon beta-1a (Rebif)	22 mcg/0.5 ml 3 times per week (IM)	Pre-filled syringe or autoinjector	7,027.32	12	91,355
Interferon beta-1a (Rebif)	44 mcg/0.5 ml 3 times per week (IM)	Pre-filled syringe or autoinjector	8,904.26	12	115,755
Interferon beta-1b (Betaferon)	250 mcg /mL every other day (SC)	Powder for injec- tion	4,937.05 (start package) 5,513.18	1	66,318
Interferon beta-1b (Extavia)	250 mcg /mL every other day (SC)	Powder for injec- tion	4,950.14	15	60,062
Natalizumab (Tysabri)	300 mg/15 mL Every four weeks (IV)	Vial	14,757.51	1	191,848
Peg-interferon beta-1a (Plegridy)	63 mcg/0.5 ml (first dose), 94 mcg/0.5 ml (second dose), 125 mcg/0.5 ml every 14 days (SC)	Prefilled syringe	8,820.69 (start package) 8,820.69	1 (63 mcg) and 1 (94 mcg) 2	114,669
Teriflunomide (Aubagio)	14 mg/day	Tablet	24,249.21	84	105,369

IM: intramuscular; IV: intravenous; mcg: microgram; mg: milligram; SC: subcutaneous ^a Source: Drug procurement cooperation (LIS) 2015.

^b The majority of patients receiving Alemtuzumab would not need new treatment after 5 year treatment. It was assumed that 20% of patients need extra treatment (12 mg/day for 3 days) (84).

^c Glatiramer acetate 40 mg/ml 3 times per week: LIS price 2015: 6702,38 (12 syringes per package). Annual drug cost was estimated to be NOK 87,131.

Monitoring costs associated with the use of medicines

Monitoring costs associated with use of medicines were calculated based on the estimates that we received from the drug procurement cooperation (LIS). The monitoring costs were estimated separately for the first and second year. Based on the information from clinical experts, we calculated the monitoring costs beyond the second year. The estimated monitoring costs are summarized in Table 23 and Appendix 8.

Drug	1. year	2. year	Beyond 2. year
Alemtuzumab ª (Lemtrada)	22,735	14,573	8307 (35.year) 7075 (+5.year)
Dimethyl fumarate (Tecifidera)	11,550	7075	7075
Fingolimod (Gilenya)	17,912	7075	7075
Glatiramer acetate (Copaxone)	11,550	7075	7075
Interferon beta-1a 30 mcg (Avonex)	19,266	14,791	7075
Interferon beta-1a 22 mcg (Rebif)	19,266	14,791	7075
Interferon beta-1a 44 mcg (Rebif)	19,266	14,791	7075
Interferon beta-1b (Betaferon)	19,266	14,791	7075
Interferon beta-1b (Extavia)	19,266	14,791	7075
Natalizumab (Tysabri)	33,240	27,725	27,725
Peg-interferon beta-1a (Plegridy)	19,266	14,791	7075
Teriflunomide (Aubagio)	12,894	7523	7523

* All costs were updated to 2015 costs.

^a The majority of patients receiving alemtuzumab would not need new treatment after 5 - year treatment. It was assumed that 20% of patients need extra treatment (12 mg/day for 3 days) (84).

Costs associated with MS care (exclusive costs associated with interventions)

The costs associated with different health states (EDSS levels) were obtained from a Norwegian study (85). This was a survey study carried out in Hordaland county in 2013 including 546 MS patients. The costs related to diagnosis, treatment, nursing care, assistive devices and equipment were included in the cost calculation.

The costs of mild or moderate and severe relapse were estimated based on the survey carried out by Svendsen in 2013 (85). The difference between the monthly costs for patients who had experienced relapse and for those who had not experienced relapse were estimated to be approximately NOK 14,600.

The cost associated to different EDSS states and relapse are presented in Table 24.

EDSS	Direct costs ^b (NOK)
0	18,046
1	36,901
2	51,297
3	126,145
4	147,554
5	329,743
6	564,928
7	689,224
8	1,380,296
9	1,393,636
Cost per relapse ^c	
Mild/ moderate	21,906
Severe	43,812

Table 24. Costs associated to different EDSS states a

EDSS: Expanded Disability Status Scale

- ^a Estimated costs associated to different EDSS states in Norway (2013) (85). All costs were updated in 2015 NOK (based on the consumer price index for the first four months of 2015 (74)).
- **b** Including VAT
- c It was assumed that the average length of mild or moderate relapse and severe relapse would be 45 and 90 days, respectively (27, 81).

We assumed that most of the patients who developed PML needed treatment at hospital. The costs were estimated based on prices from the Norwegian DRG system (DRG code 421; personal communication by dr.med Elisabeth Gulowsen Celius). Patients who survived PML also needed 3-6 months extra treatments at rehabilitation centres. We assumed NOK 3,000 cost per day for stay at rehabilitation centre (86).

Health-related Quality of Life

In order to obtain utility weights, we performed a systematic search for published values. For consistency, and as the use of different utility instruments would yield different results, we focused on values based on EQ-5D, the most commonly used instrument (87).

In the base-case, we used the utility values reported by Orme and co-workers (88). The study was a cross sectional study of people comprising all course of MS (RRMS, SPMS and PPMS) from the United Kingdom. Based on the systematic search on health related quality of life data, this is the only study that presented the utility associated with each EDSS state, SPMS and relapse by using the EQ-5D method.

As Orme and colleagues did not make a distinction between mild or moderate and severe relapse, we assumed that the reported disutility was for mild or moderate relapses. Therefore, the ratio between disutility associated with mild or moderate relapse and severe relapse estimated by Prosser and co-workers (81) was applied to estimate the disutility associated with severe relapse. As mentioned, it was assumed that the average length of mild or moderate relapse and severe relapse would be 45 and 90 days, respectively (27, 81).

We assumed a disutility of 0.4 (0.3-0.5) assigned to the year a patient experienced PML (89).

Beta or log-normal distributions were used for utility values used in the model. The mean values and standard errors of the utility (QALY) weights used in our model are presented in the Table 25.

We did not identify reliable data on the probable effect on patients' utility of the different methods of administrating the medication. Therefore, the possible disutility associated with injections is not included in the model.

Table 25. Quality of life data (base-case)

Parameter	Utility weight	95% CL		Probability distribution
EDSS 0	0.870	0.782	0.958	Beta
EDSS 1	0.799	0.799	0.617	Beta
EDSS 2	0.705	0.705	.0523	Beta
EDSS 3	0.574	0.574	0.384	Beta
EDSS 4	0.610	0.610	0.428	Beta
EDSS 5	0.518	0.518	0.338	Beta
EDSS 6	0.460	0.277	0.641	Beta
EDSS 7	0.297	0.112	0.481	Beta
EDSS 8	-0.049	-0.235	-0.138	Log-normal
EDSS 9	-0.195	-0.428	-0.039	Log-normal
SPMS ^a	-0.045	-0.076	-0.014	Beta or Log-normal
Disutility associated with mild or moderate relapse	-0.071	-0.096	-0.046	Log-normal
Disutility associated with severe relapse ^b	-0.236	-0.295	-0.174	Log-normal
Disability associated with PML °	-0.40	-0.30	-0.50	Log-normal

CI: confidence interval; EDSS: Expanded Disability Status Scale; SPMS: Secondary Progressive MS ^a Assumed fixed utility decrement over the corresponding RRMS EDSS state utility values. ^b It was estimated based on the data reported by Orme et al. (88) and Prosser et al. (81).

c Ref:(89)

Economic evaluation – Results

We calculated costs and effectiveness (in terms of QALYs), for all relevant disease modifying therapies used for RRMS based on simulations of the model. We used 10,000 iterations in the Monte Carlo analyses. Our assessment of cost-effectiveness will reflect a range of potential willingness to pay (WTP) values per gained QALY.

Incremental cost-effectiveness estimates

The results of the base-case analysis are presented in Table 26. Over a 20-year time horizon, alemtuzumab dominated all other alternative treatments, *i.e.* it was both more effective and less costly.

Drugs	Total costs (NOK)	Effects (QALYs)	Incremental cost (NOK)	Incremental effect (QALYs)	ICER (NOK/QALY)
Alemtuzumab (Lemtrada)	4,897,903	8.05			Dominant
Interferon beta-1b (Extavia)	6,031,551	7.40	1,133,647	-0.64	Dominated by alemtuzumab
Interferon beta-1b (Betaferon)	6,088,153	7.40	1,190,250	-0.64	Dominated by alemtuzumab
Glatiramer acetate 20mg (Copax- one)*	6,253,728	7.31	1,355,825	-0.73	Dominated by alemtuzumab
Peg-interferon beta-1a (Plegridy)	6,310,586	7.56	1,412,682	-0.48	Dominated by alemtuzumab
Teriflunomide (Aubagio)	6,337,489	7.38	1,439,586	-0.67	Dominated by alemtuzumab
Interferon beta-1a 22 mcg (Rebif)	6,498,571	7.21	1,600,667	-0.84	Dominated by alemtuzumab

Table 26: Results of the base-case cost-effectiveness analysis (discounted)

Interferon beta-1a 30 mcg (Avonex)	6,533,915	7.27	1,636,012	-0.77	Dominated by alemtuzumab
Interferon beta-1a 44 mcg (Rebif)	6,574,606	7.32	1,676,702	-0.72	Dominated by alemtuzumab
Dimethyl fumarate (Tecifidera)	6,707,787	7.52	1,809,884	-0.52	Dominated by alemtuzumab
Natalizumab (Tysabri)	6,983,132	7.63	2,085,228	-0.41	Dominated by alemtuzumab
Fingolimod (Gilenya)	7,041,216	7.43	2,143,313	-0.62	Dominated by alemtuzumab

QALY: quality-adjusted life year; ICER: incremental cost-effectiveness ratio * Based on effect estimates and annual drug costs, it is highly probable that glatiramer acetate 40 mg 3 times per week will be as cost-effective as glatiramer acetate 20 mg per day (given all the other parameters are the same).

Monte Carlo simulations with 10,000 draws from the input distributions are shown in Figure 10. Simulations for alemtuzumab show that alemtuzumab was more effective and less costly relative to other treatments. All other interventions were dominated by alemtuzumab. The results of the probabilistic sensitivity analysis also showed that alemtuzumab was more likely to be the most cost-effective strategy (above 90%) for all values of WTP.

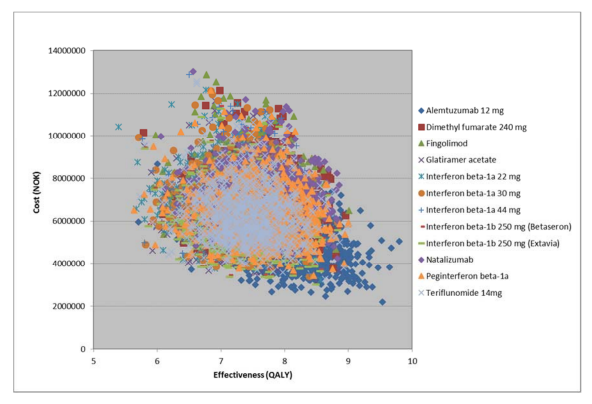


Figure 10. Cost-effectiveness scatter-plot; mcg: microgram; mg: milligram

The results presented above show that alemtuzumab was the most cost-effective strategy and dominated all other treatment strategies. In order to show the cost-effectiveness of other treatment strategies relative to each other, we excluded alemtuzumab (the dominate strategy) and conducted a separate analysis of the remaining interventions. The results (for all treatment strategies, except alemtuzumab) are presented in Table 27 and Figure 11.

Discarding alemtuzumab, natalizumab was the most effective treatment regarding QALYs (7.63), followed by peg-interferon beta-1a (7.56). Interferon beta-1a 22 mcg was the least effective strategy (7.21).

Fingolimod was the most expensive treatment (NOK 7,050,000), followed by natalizumab (NOK 6,984,840). Interferon beta-1b (Extavia) was the least expensive treament (NOK 6,033,330) and was, therefore, used as a reference.

Three treatment strategies were not dominated by the other interventions. The incremental cost per QALY for peg-interferon beta-1a versus interferon beta-1b (Extavia) was NOK 1,658,450. The incremental cost per QALY for natalizumab versus peg-interferon beta-1a was NOK 10,620,000.

Interferon beta-1b (Betaferon) was dominated by interferon-1b (Extavia); glatiramer acetate was dominated by interferon beta-1b (Extavia and Betaferon), while teriflunomide was dominated by interferon beta-1b (Extavia and Betaferon) and peg-interferon beta-1a.

Interferon beta-1a (Rebif and Avonex) was dominated by peg-interferon beta-1a, interferon beta-1b (Extavia and Betaferon), teriflunomide, and glatiramer acetate. Dimethyl fumarate was dominated by peg-interferon beta-1a, while fingolimod was dominated by natalizumab, peg-interferon beta-1a and dimethyl fumarate.

Table 27. Results of the base-case cost-effectiveness analysis (all interventions except
alemtuzumab) (discounted)

Drugs	Total costs	Effects	Versus Inte	rferon beta-1b 250	Sequential ICER		
2.490	(NOK)	(QALYs)	Incremental cost (NOK)	Incremental effect (QALYs)	ICER (NOK/QALY)	(NOK/QALY)	
Interferon beta-1b (Extavia)	6,033,328	7.40					
Peg-interferon beta-1a (Plegridy)	6,308,924	7.56	275,597	0.17	1,658,451	1,658,451	
Natalizumab (Tysabri)	6,984,843	7.63	951,515	0.23	4,140,203	10,619,960	

Dominated therap	ies					
Interferon beta-1b (Beta- feron)	6,089,587	7.40	56,259	-	Dominated by interferon beta- 1b (Extavia)	Dominated by interferon beta-1b (Extavia)
Glatiramer ace- tate 20 mg (Co- paxone) *	6,256,047	7.31	222,720	-0.09	Dominated	Dominated by interferon beta-1b (Extavia) and in- terferon beta-1b (Beta- feron)
Teriflunomide (Aubagio)	6,332,443	7.38	299,116	-0.02	Dominated	Dominated by interferon beta-1b (Extavia), inter- feron beta-1b (Betaferon) and peg-interferon beta- 1a
Interferon beta-1a 22 mcg (Rebif)	6,497,728	7.21	464,401	-0.19	Dominated	Dominated by interferon beta-1b (Extavia), inter- feron beta-1b (Betaferon), peg-interferon beta-1a, glatiramer acetate and teriflunomide
Interferon beta- 1a 30 mcg (Avonex)	6,539,464	7.27	506,137	-0.13	Dominated	Dominated by interferon beta-1b (Extavia), inter- feron beta-1b (Beta- feron), peg-interferon beta-1a, glatiramer ace- tate and teriflunomide
Interferon beta-1a 44 mcg (Rebif)	6,573,653	7.32	540,325	-0.08	Dominated	Dominated by interferon beta-1b (Extavia), inter- feron beta-1b (Betaferon), peg-interferon beta-1a, glatiramer acetate and teriflunomide
Dimethyl fumarate (Tecifidera)	6,710,845	7.52	677,517	0.12	5,746,659	Dominated peg-interferon beta-1a
Fingolimod (Gilenya)	7,040,995	7.42	1,007,668	0.02	43,827,412	Dominated by peg-inter- feron beta-1a, dimethyl fumarate and natalizumab

QALY: quality-adjusted life year; ICER: incremental cost-effectiveness ratio; mcg: mi-crogram; mg: milligram * Based on effect estimates and annual drug costs, it is highly probable that glatiramer ace-tate 40 mg 3 times per week will be as cost-effective as glatiramer acetate 20 mg per day (given all the other parameters are the same).

The incremental cost versus incremental effectiveness (QALY), when all treatment strategies, except alemtuzumab are included in the analysis, is presented in Figure 11. As mentioned, three interventions, interferon beta-1b (Extavia), peg-interferon beta-1b (Extavia) to peg-interferon beta-1a and to natalizumab represent the cost-effectiveness frontier. It means that at different WTP, these three strategies could be considered the most cost-effective. The incremental cost per QALY of peg-interferon beta-1a compared with interferon beta-1b (Extavia) is estimated to be NOK 1,658,000, meaning interferon beta-1b (Extavia) could be considered the cost-effective treatment if WTP for QALY is less than NOK 1,658,000. For WTP between NOK 1,658,000 and NOK 10,620,000, peg-interferon beta-1a is the cost-effective treatment. If WTP is above 10,620,000, then natalizumab is the cost-effective treatment. The other treatments were dominated by the treatment comprising in the frontier. Therefore, they were not considered to be cost-effective.

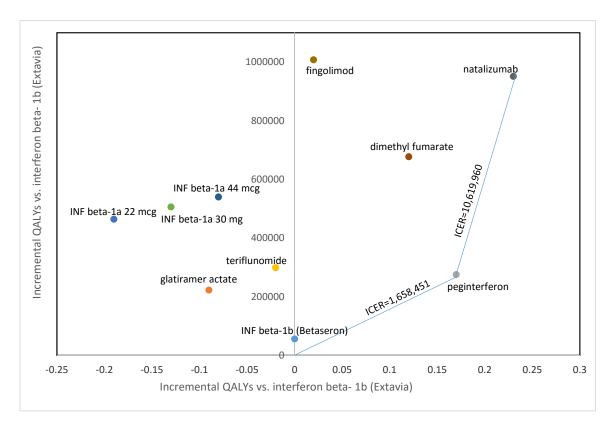


Figure 11. Cost-effectiveness graph (all interventions except alemtuzumab); mcg: microgram; mg: milligram; INf: interferon

We performed a Monte Carlo simulation with 10,000 draws from the input distributions and we varied the WTP from NOK 0 to NOK 2,000,000. The cost-effectiveness acceptability curves in Figure 12 show the probability of the alternatives being costeffective subject to different levels of WTP. If one assumes maximum WTP per QALY is NOK 500,000, interferon beta-1a (Extavia) was the most cost-effective treatment strategy (47%), followed by peg-interferon beta-1a (27%) and teriflunomide (13%). With a WPT per QALY of NOK 1,000,000, interferon beta-1b (Extavia) was the most cost-effective (36%) followed by peg-interferon beta-1a (34%) and teriflunomide (14%). However, as presented in the cost-effectiveness scatterplot (Figure 10) and Table 27, total QALYs of included interventions (except alemtuzumab) overlapped, which indicates the uncertainty regarding the gain in QALYs.

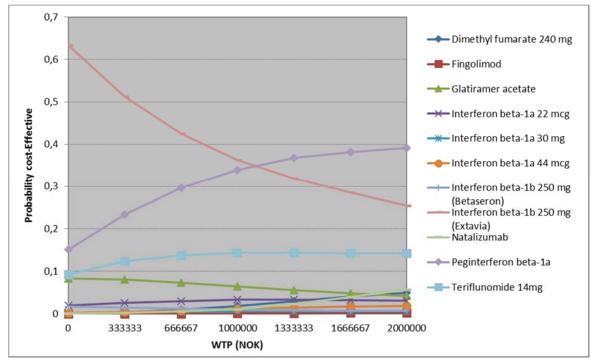


Figure 12. Cost-effectiveness acceptability curve (all interventions except alemtuzumab) WTP willingness to pay; mcg: microgram; mg: milligram

Value of information analysis

We performed an analysis of the expected value of perfect information (EVPI) on all uncertain parameters to explore the uncertainty surrounding specific groups of parameters and show which group has the most impact on the results. EVPI analyses were performed with 100x500 iterations. The EVPI of different groups of parameters (costs, efficacy, QALYs and probabilities) are presented in Figure 13.

At a WTP of NOK 400,000 per QALY, probabilities data (Norwegian epidemiological data) had the highest EVPI. For values of WTP above NOK 1,000,000 per QALY, the results indicate that the treatment efficacy data have the greatest impact on decision uncertainty. These results suggest that if new research is to be undertaken (for WTP above NOK 1,000,000), additional information on efficacy data would contribute most to reducing the uncertainty surrounding the decision about which treatment modality is most cost-effective.

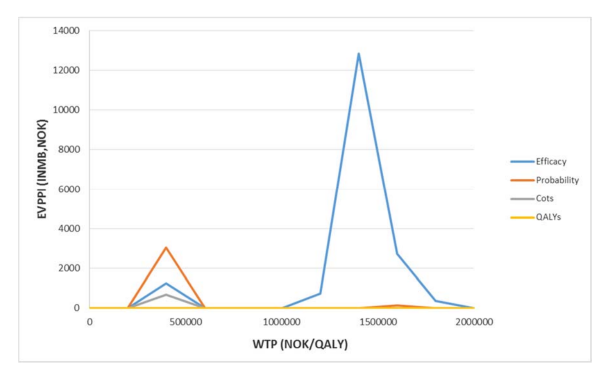


Figure 13. Expected value of partial perfect information per patient for different groups of parameters; QALY: quality-adjusted life year; WTP: willingness to pay; INMB: incremental net monetary benefit

Scenario analyses

In addition to the probabilistic sensitivity analysis, we performed several scenario analyses to test the uncertainty around the model assumptions and some of the input parameters.

"No treatment" was our common comparator in the network meta-analyses, and therefore was included in the health economics model. As additional information, we presented the cost-effectiveness of all treatment strategies compared to "no treatment" as a scenario analysis. The results are presented in Table 28. They showed that alemtuzumab remained the dominant strategy (less costly and more effective). Interferon beta-1b (Extavia and Betaferon) had ICERs below NOK 500,000 per QALY. Peg-interferon beta-1a had ICER between NOK 500,000-800,000 per QALY. Teriflino-mide and glatiramer acetate had ICERs between NOK 1,000,000-1,500,000 per QALY. Dimethyl fumarate and natalizumab had ICERs between 1,500,000-1,800,000 per QALY. Interferon beta-1a (22mcg, 44 mcg and 30 mcg) and fingolimod had ICERs above NOK 2,000,000 per QALY.

Table 28. Cost-effectiveness of disease-modifying therapies compared to "no treatment" (discounted)

Drugs	Total costs (NOK)	Effects (QALYs)	Incremental cost (NOK)	Incremental effect (QALYs)	ICER (NOK/QALY)
No treatment	5,900,815	7.00			
Alemtuzumab (Lemtrada)	4,897,903	8.05	-1,002,911	1.05	Dominant
Interferon beta-1b (Extavia)	6,031,551	7.40	130,736	0.40	326,841
Interferon beta-1b (Beta- feron)	6,088,153	7.40	187,339	0.40	468,346
Glatiramer acetate (Copaxone) *	6,253,728	7.31	352,914	0.31	1,138,431
Peg-interferon beta-1a (Plegridy)	6,310,586	7.56	409,771	0-56	731,734
Teriflunomide (Aubagio)	6,337,489	7.38	436,675	0.38	1,149,144
Interferon beta-1a 22 mcg (Rebif)	6,498,571	7.21	597,756	0.21	2,846,458
Interferon beta-1a 30 mcg (Avonex)	6,533,915	7.27	633,101	0.27	2,344,817
Interferon beta-1a 44 mcg (Rebif)	6,574,606	7.32	673,791	0.32	2,105,598
Dimethyl fumarate (Tecifidera)	6,707,787	7.52	806,973	0.52	1,551,870
Natalizumab (Tysabri)	6,983,132	7.63	1,082,317	0.63	1,717,964
Fingolimod (Gilenya)	7,041,216	7.43	1,140,402	0.43	2,652,097

QALY: quality-adjusted life year; ICER: incremental cost-effectiveness ratio

* Based on effect estimates and annual drug costs, it is highly probable that glatiramer acetate 40 mg 3 times per week will be as cost-effective as glatiramer acetate 20 mg per day (given all the other parameters are the same).

In the base-case analysis, we assumed that once patients progress to EDSS=6 or SPMS, they would not receive MS treatment anymore. A scenario analysis was conducted varying the EDSS levels where treatment would be discontinued. The results of scenario analysis showed that ICERs were reduced when considering a stopping rule at EDSS=7 (Appendix 9.1). We also assumed a stopping rule without considering SPMS progression. As we did not consider any treatment benefit for SPMS patients in our model, a scenario analysis without considering treatment discontinuation with the progression to SPMS resulted in much higher ICERs.

A time horizon of 20 years was considered in the base-case analysis. We performed a scenario analysis where the time horizon varied within the range of 10 years. A time horizon of 30 years resulted in lower ICERs (Appendix 9.2), and the scenario analysis indicated that a time horizon of 10 years would increase the ICERs.

We also conducted a scenario analysis where the starting age was changed within the range of 10 years. Scenario analysis showed that variation in the starting age had a very small potential impact on the results. However, treating younger patients would slightly decrease the ICERs.

For base-case analysis, we assumed disability improvements (a maximum of 2 EDSSlevel). We performed a scenario analysis where no improvement in EDSS were modelled. ICERs were not very sensitive to this assumption. However, "no improvement" in EDSS-level resulted in slightly lower ICERs (Appendix 9.3).

The annual rate of treatment discontinuation was assumed to be 15% in the base-case analysis. Based on our systematic review the rate varied between 0 and 33%. We conducted two scenario-analyses where the annual rate of treatment discontinuation was considered to be 0 and 30%, respectively. The scenario analyses showed that discontinuation rate did not have a significant impact on the results.

Utility values reported by Orme and co-workers (88) were used in the base-case analysis, as it was the only study that presented the utility associated with EDSS-states, SPMS and relapse by using a generic preference-based instrument (EQ-5D). We performed a scenario analysis based on utility values reported by Svendsen and coworker (90). Utility values were calculated based on data from 423 Norwegian patients by using the EQ-5D method (Table 29).

Table 29. Quality life data reported by Svendsen et al. (90)

	EDSS 1	EDSS 2	EDSS 3	EDSS 4	EDSS 5	EDSS 6	EDSS 7	EDSS 8	EDSS 9
Quality of life	0.800	0.757	0.701	0.617	0.536	0.443	0.211	0.142	0.056

The use of different quality of life data resulted in different QALYs gained (higher QALYs for all interventions). However, the conclusion remained the same as in the base-case analysis. The results are presented in Appendix 9.4.

It has been reported that more patients (about 22-28%) than we assumed may need three cycles of alemtuzumab during the 5-year period (and some patients may need four (about 8-10% of patients) or five cycles (1.5%) of alemtuzumab). The scenario analysis was performed by varying the probability of patients who need more than 2 cycles of alemtuzumab. The results showed that alemtuzumab still was the dominant strategy.

Budget impact

The prevalence of MS in Norway is estimated to be 203 per 100,000 people (8). Approximately 85%-90% of patients with MS are estimated to have RRMS from onset of disease (11). We assumed that about 50% of these patients are eligible for disease modifying therapies, based on a Norwegian study (91). Based on these assumptions, we have estimated the number of eligible patients for disease-modifying therapies for the next 5 years (Table 30).

	2015 *	2016	2017	2018	2019	2020
Number of patients	4610	4650	4690	4740	4780	4830

*The population used in the analysis was 5,165,802 which was the population in Norway in 1. January 2015. It was assumed that the population of Norway increases about 50,000 annually (74) The market shares for disease-modifying therapies for the last three years is presented in Figure 14 and Table 31, based on sales data (defined daily dose; DDD) (Farmastat). As results show, in the past few years the oral MS-medicines won market share from non-oral treatment alternatives.

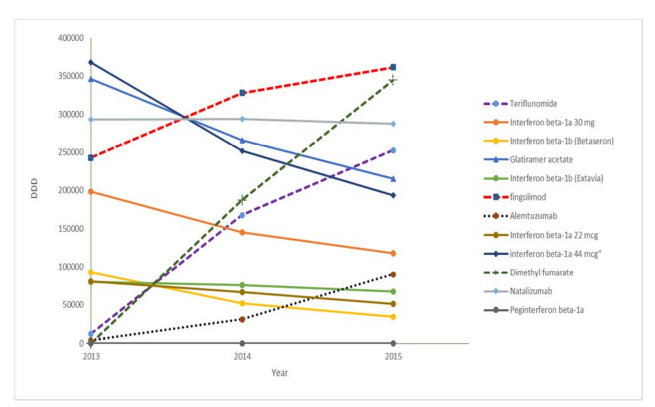


Figure 14. Sales data for disease-modifying therapies in DDD (Farmastat) DDD: defined daily dose; Sales data for 2015 were estimated based on data from the first half of 2015.

Drugs	2013	2014	2015 ª
Alemtuzumab (Lemtrada)	0%	2%	4%
Dimethyl fumarate (Tecifidera)	0 %	10%	17%
Fingolimod (Gilenya)	14%	18%	18%
Glatiramer acetate (Copaxone)	20%	14%	11%
Interferon beta-1a 30 mcg (Avonex)	12%	8%	6%
Interferon beta-1a 22 mcg (Rebif)	5%	4%	3%
Interferon beta-1a 44 mcg (Rebif)	21%	14%	10%
Interferon beta-1b (Betaferon)	5%	3%	2%
Interferon beta-1b (Extavia)	5%	4%	3%
Natalizumab (Tysabri)	17%	16%	14%
Peg-interferon beta-1a (Plegridy) ^b	0%	0%	0%
Teriflunomide (Aubagio)	1%	9%	13%

Table 31. Current market shares for disease-modifying therapies in DDD (Farmastat)

DDD: defined daily dose

^a Estimated based on data from the first half of 2015.

^b Peg-interfron beta-1a: DDD 2013=0, DDD 2014= 70, DDD 2015=337

The market share forecasts for the next five years were estimated based on the results of our cost-effectiveness analysis and the drugs' adverse events. We also took under consideration the current practice where there is a trend in favour of oral medicines. The results were presented in Table 32.

Drugs	2016	2017	2018	2019	2020
Alemtuzumab (Lemtrada)	15%	19%	24%	31%	33%
Dimethyl fumarate (Tecifidera)	13%	13%	12%	11%	10%
Fingolimod (Gilenya)	13%	12.5%	12%	12%	12%
Glatiramer acetate (Copaxone)	7%	6%	5%	4%	3%
Interferon beta-1a 30 mcg (Avonex)	4%	3%	2%	1%	1%
Interferon beta-1a 22 mcg (Rebif)	2%	1.5%	1%	0%	0%
Interferon beta-1a 44 mcg (Rebif)	8%	5%	3%	1%	1%
Interferon beta-1b (Betaferon)	2%	1%	1%	0%	0%
Interferon beta-1b (Extavia)	9%	9%	9%	9%	9%
Natalizumab (Tysabri)	12%	12%	12%	12%	12%
Peg-interferon beta-1a (Plegridy)	4%	4%	4%	4%	4%
Teriflunomide (Aubagio)	14%	14%	15%	15%	15%

Table 32: Forecasted marked shares for disease-modifying therapies

The budget impact was calculated based on the same cost inputs (drug costs, monitoring costs associated with use of drugs) used in the cost-effectiveness model (see Tables 22 and 23). All estimations are based on 2015-price. The results of the budget impact analysis for the next five years (2016 was assumed as a starting point) are shown in Tables 33-35. Table 33 presented estimated costs based on current practice, while Table 34 presented estimated costs based on future practice (based on data from Table 32). Estimated costs based on future practice compared to estimated costs based on current practice were presented in Table 35.

Table 33. Estimated costs* based on current practice

Drugs	2016	2017	2018	2019	2020
Alemtuzumab (Lemtrada)	70,957,237	43,384,319	5,405,710	5,381,873	5,488,448
Dimethyl fumarate (Tecifidera)	143,409 155	140,972,076	142,482,533	143,676,691	145,187,149
Fingolimod (Gilenya)	179,884,866	172,095,248	173,949,176	175,396,174	177,250,102
Glatiramer acetate (Copaxone)	49,276,655	47,406,610	47,916,734	48,315,953	48,826,077
Interferon beta-1a 30 mcg (Avonex)	33,691,593	32,727,302	30,916,393	31,174,127	31,502,362
Interferon beta-1a 22 mcg (Rebif)	13,167,637	12,733,796	11,927,325	12,026,598	12,153,335
Interferon beta-1a 44 mcg (Rebif)	60,609,419	59,065,448	56,138,239	56,606,768	57,202,424
Interferon beta-1b (Beta- feron)	6,899,458	6,588,894	6,022,076	6,071,961	6,136,109
Interferon beta-1b (Extavia)	12,465,793	11,852,383	10,737,660	10,826,455	10,940,936
Natalizumab (Tysabri)	149,923 462	147,436,954	149,016,551	150,265,611	151,,845,208
Peg-interferon beta-1a (Plegridy)	104,602	101,908	96,806	97,614	98,642
Teriflunomide (Aubagio)	69,119,685	66,483,177	67,198,605	67,758,440	68,473,868
Total	789,509,563	740,848,115	701,807,807	707 598 265	715,104,659

* Undiscounted costs, included VAT

Table 34. Estimated costs* based on future practice

Drugs	2016	2017	2018	2019	2020
Alemtuzumab (Lemtrada)	190,987,053	184,889,553	29,099,797	37,421,494	40,624,633
Dimethyl fumarate (Tecifidera)	109,173,395	107,318,114	100,124,293	92,549,822	85,020,717
Fingolimod (Gilenya)	130,374,387	119,931,479	116,374,524	117,342,587	118,582,892
Glatiramer acetate (Copaxone)	32,322,524	26,653,617	22,450,355	18,109,921	13,725,845
Interferon beta-1a 30 mcg (Avonex)	23,098,927	16,828,357	10,598,126	5,343,239	5,429,044
Interferon beta-1a 22 mcg (Rebif)	10,346,880	7,504,483	4,686,134	0	0
Interferon beta-1a 44 mcg (Rebif)	50,464,321	30,736,743	17,528,081	5,891,457	5,953,451
Interferon beta-1b (Beta- feron)	8,018,439	3,828,753	3,499,380	0	0
Interferon beta-1b (Extavia)	33,464,841	31,818,121	28,832,186	29,063,982	29,371,310
Natalizumab (Tysabri)	126,211,185	124,112,685	125,447,714	126,499,219	127,828,983
Peg-interferon beta-1a (Plegridy)	25,030,165	24,385,660	23,164,869	23,358,184	23,603,988
Teriflunomide (Aubagio)	77,266,215	74,318,964	80,484,336	81,154,855	82,011,729
Total	816,758,333	752,326,530	562,289,795	536,734,760	532,152,591

* Undiscounted costs, included VAT

Table 35. The results of the budget impact; estimated costs based on future practice compared to estimated costs based on current practice

Drugs	2016	2017	2018	2019	2020
Alemtuzumab (Lemtrada)	120,029,816	141,505,234	23,694,087	32,039,621	35,136,185
Dimethyl fumarate (Tecifidera)	-34,235,760	-33,653,961	-42,358,241	-51,126,869	-60,166,432
Fingolimod (Gilenya)	-49,510,480	-52,163,769	-57,574,651	-58,053,587	-58,667,210
Glatiramer acetate (Copaxone)	-16,954,131	-20,752,993	-25,466,378	-30,206,032	-35,100,232
Interferon beta-1a 30 mcg (Avonex)	-10,592,666	-15,898,945	-20,318,267	-25,830,888	-26,073,319
Interferon beta-1a 22 mcg (Rebif)	-2,820,756	-5,229,314	-7,241,192	-12,026,598	-12,153,335
Interferon beta-1a 44 mcg (Rebif)	-10,145,097	-28,328,706	-38,610,158	-50,715,312	-51,248,973
Interferon beta-1b (Beta- feron)	1,118,981	-2,760,140	-2,522,696	-6,071,961	-6,136,109
Interferon beta-1b (Extavia)	20,999,048	19,965,738	18,094,526	18,237,527	18,430,374
Natalizumab (Tysabri)	-23,712,276	-23,324,269	-23,568,837	-23,766,392	-24,016,225
Peg-interferon beta-1a (Plegridy)	24,925,563	24,283,752	23,068,062	23,260,569	23,505,346
Teriflunomide (Aubagio)	8,146,530	7,835,788	13,285,731	13,396,415	13,537,861
Total	27,248,771	11,478,415	-139,518,013	-170,863,506	-182,952,068

The budgetary impact for the next 5 years is difficult to predict. The prediction depends on several factors, including any change in current clinical practice, the relative drug prices and the number of patients eligible for different treatment alternatives.

For budget impact analysis, we mainly assumed that alemtuzumab, the more effective and less costly treatment alternative, would capture higher market share in the future. The results presented in Table 35 showed that in the first two years, there will be additional costs compared to costs estimated based on current practice. However, our results indicated that costs would decrease after the first two years and there is a potential for cost-savings. Overall, the potential cost-savings over a 5-year period were estimated to be NOK 454,606,000 compared to the costs estimated for current practice.

Discussion

In this HTA, we have systematically reviewed the literature on the clinical effect of disease modifying medicines used for multiple sclerosis. The evidence base comprised findings from 37 RCTs. Furthermore, we performed an economic evaluation to examine the cost-effectiveness of these disease-modifying medicines in a Norwegian setting.

Summary of key findings

Key findings of the clinical evaluation

All examined treatments were more effective than placebo against annual relapse. The strongest effect was seen for alemtuzumab 12 mg. Fingolimod oral 1.25 mg and dimethyl fumarate 240 mg two times a day were also associated with a reduction in annualised relapse rate.

For disability progression, there is high quality evidence showing that dimethyl fumarate 240 mg twice daily and fingolimod oral 0.5 mg are more effective than placebo. For withdrawal due to adverse events, the lower quality of the available evidence provides unclear conclusion. Results indicate that some treatments are associated with more withdrawal due to adverse events than placebo, such as interferon beta-1a 44 mcg, and all regimens of peg-interferon beta-1a mcg.

For change in disability status, serious adverse events and mortality, we did not access the quality of the available evidence. Therefore, one cannot conclude on how reliable results are for these outcomes. Our results indicate that interferon beta-1a 30 mcg is related to a negative progression in disability status scale. Finally, our results did not show that examined treatments increased mortality.

Key findings of economic evaluation

Our health economic analysis indicated that alemtuzumab was more effective and less costly than the other treatment alternatives dominating all other disease-modifying therapies.

A scenario analysis that excluded alemtuzumab (the dominant strategy) showed that natalizumab was the most effective (in terms of QALYs), and interferon beta-1a 22 mg was the least effective treatment. Fingolimod was the most expensive strategy and interferon beta-1b was the least expensive alternative. The results also showed that only three treatment alternatives (interferon beta-1b (Extavia), peg-interferon beta-1a and natalizumab) could be cost-effective depending on the willingness-to-pay (WTP) threshold. Interferon beta-1b was likely to be the cost-effective choice for a WTP per QALY below NOK 1,658,000. Peg-interferon was the cost-effective option for a WTP from NOK 1,658,450 to NOK 10,619,960, and natalizumab was the cost-effective alternative for a WTP above NOK 10,619,960. Assuming a WTP below NOK 1,000,000 per QALY, interferon beta-1b (Extavia) was approximately 40% likely to be the most cost-effective treatment, followed by peg-interferon beta-1a (approximately 30% likely).

The scenario analysis where all treatment alternatives were compared to "no treatment" indicated that alemtuzumab remained the dominant strategy. Interferon beta-1b had ICERs below NOK 500,000 per QALY. The ICER for peg-interferon compared to "no treatment" was NOK 731,730. Other treatment options had ICERs over NOK 1,000,000 per QALY. The treatment costs (included drug costs and monitoring costs associated with each treatment) had an impact on the ICERs.

The results of probabilistic analysis showed that there is some degree of uncertainty regarding the input parameters. More research on efficacy and epidemiologic input parameters would have the greatest impact on reducing decision uncertainty.

In addition to our probabilistic sensitivity analysis, we performed several scenario analyses to test the uncertainty around the model assumptions. The results showed that, while there were numerical changes to the ICERs, the cost-effectiveness results were robust to variations in the model assumptions and the conclusions of the analysis would not change.

Our bugdet impact analysis based on the results of our cost-effectiveness analysis, the drugs' adverse events profile, and current clinical practice showed that there is a substantial potential for cost saving.

Quality of the evidence

Quality of the evidence of the systematic review

We included a HTA of high quality. We updated the information with more recently published RCTs with generally low risk of bias.

We chose a conservative approach in grading the quality of the evidence. This implies that one can rely on the evidence we judged to be of high quality. We had evidence of high quality only for annual relapse rates and disability progression. This implies that results on other outcomes are less reliable.

Quality of the economic evaluation

Our cost-effectiveness analysis showed that there is some degree of uncertainty around the estimates. This was mainly due to uncertainty in the efficacy data, followed by probabilities estimates.

Strengths and weaknesses

Strengths of the systematic review

We used an internationally recognised methodology to systematically search the evidence, extract the data, access bias of studies and the quality of evidence. While the focus of this report was MS treatments used in Norway, we included evidence for treatments that are both used in Norway and not to get a bigger network of evidence for medicines relevant to the Norwegian setting. Our network of evidence includes information on treatments that have been used for some years, and on emerging treatments.

Limitations of the systematic review

Many of the limitations of this report are related to the available evidence, and are not inherent to the methodology used in this report.

The available evidence differs by treatments according to how long these have been on the market, with newer treatments having a smaller amount of information.

Most MS medications are only approved for RRMS patients. The systematic review includes, therefore, only studies of RRMS patients. As RRMS patients represent the largest proportion of MS patients, the results of our report are relevant to the majority of MS patients in Norway. Furthermore, there is no reason to believe that the effect of these medications are different depending on if one treats after the first relapse (CIS scenario) or if the treatment is initiated after the second relapse (e.g. definite clinical MS including RRMS patients). Results related to newer medications carry more uncertainty. As MS diagnosis has changed through the years, studies conducted at a different time might differ in terms of the MS population included. Therefore, when comparing older with newer MS treatments, differences in results could partly be due to differences in patient population. Furthermore, follow-up time of newer medicines is usually shorter, and some serious adverse events might only occur after a longer use of the medicine. One should bear this in mind when interpreting results.

Through network meta-analysis, one can infer on the relationship between two treatments if those treatments were compared to a common comparator in RCTs. For such an inference to be accurate, the contributing RCTs should be very similar regarding patient population and outcome definition, measurement and reporting. Treatment history among patients varied across the trials, being either unclear, treatment naive, treatment experienced or a mixture. However, different statistical analyses provided similar results, and results were consistent when considering direct evidence, indirect evidence from the whole network.

The available evidence does not allow to investigate separately first and second line treatments. Most published studies did not examine first and second medications separately. Indeed, some studies have compared first and second line treatments. Furthermore, in some case, first-line treatments have been investigated in patients who had taken other medications before, hence considered as second line treatments. Finally, studies considered second-line treatments in a population that comprised patients who had not received any treatment before, and were therefore tested as first-line treatments. We, therefore, present results for all MS treatments together (independent of them being used as first or second line treatments). However, patients who use a first and a second treatment might differ, and discrepancies in treatments efficacy might be due to disparity in patients.

The clinical endpoints covered in the systematic review (clinical relapse and disability progression) are important clinical outcomes in MS. Magnetic resonance imaging (MRI) is a surrogate endpoint and, therefore, was not examined. However, a previous published HTA report described that the available evidence on MRI was of poorer quality compared to clinical relapse and disability progression (27). The population of studies examining MRI populations were usually smaller, and it is unclear how these populations were selected (27). Therefore, any conclusions on MS medicines use on that surrogate outcome would have a higher degree of uncertainty.

Some outcome definitions differed from one study to the other. For example, disability progression was measured as disability progression confirmed at 3 months, or confirmed at 6 months, or at two years, or as a change compared to baseline EDSS. Patients EDSS classification might also differ between studies.

The lengths of the included studies were relatively short with a maximum follow-up time of 3.5 years. Therefore, our results cannot conclude on the long-term effect of examined medicines. Observational follow-up studies, with a longer follow-up time have been published, and could be used to estimate the longer-term effect of MS medicines.

All these limitations would not only have an impact on the clinical effect results but could also influence the health economic evaluation results that incorporated some of the clinical effect results into the health economic model.

Strengths of the health economic model

We performed the economic evaluation of disease- modifying therapies based on a thorough systematic review of the literature, and estimates of treatment effect obtained through a network meta-analysis. We used a probabilistic Markov-model, considered the appropriate approach for simulating the natural history of multiple sclerosis. This model was previously used in a high quality HTA report. The model structure and all assumptions have been adapted to the Norwegian setting based on Norwegian clinical practice with close assistance of experts in this field.

Limitations of the health economic model

To model real life is very complex; hence, any simulation is a simplification. We have tried to find the most robust and best evidence available but limitations associated with the data, and the simplifications of our health economic model should be considered when interpreting the results.

Data from Norwegian MS-registry or Norwegian cohort studies should ideally be used in the model. However, we were not able to identify data sources that were compatible to the developed model. The transitional probabilities were therefore based on estimates reported in the published literature. Those were also used in previous health economic studies. Data on annual relapse rate were uncertain. Indeed, we were not able to identify any study that linked rates of annual relapse to different EDSS-scores by disease duration.

We found a Norwegian study from 1996 where EDSS distributions in the cohort patients were reported (92). 22.6% of the patients in this study had EDSS scores over 4.5 (6.4% of patients scored between 8 and 9.5). However, based on clinical experts' opinion, disease-modifying therapies are usually initiated in patients with an EDSS score less than 5, and most commonly for patients with an EDSS score between 1 and 3. Therefore, EDSS distributions used in our model were based on published literature of large cohort studies where over 91% of patients had EDSS scores less than 5.

The network meta-analyses were not performed separately for first and second line treatments. Therefore, we did not perform separate cost-effectiveness analyses for these two types of treatments. In addition, based on expert opinion, we did not include combination therapy in our model, as it is not relevant to Norwegian clinical practice at present.

There is lack of documentation regarding the long-term effect of the newer drugs. Further research could change current estimates and consequently the health economic results. In our report, we assumed that 20% of patients might need three cycles of alemtuzumab during a 5-year period. However, it has been reported that this proportion might be higher (22 to 28% of patients), and that some patients may need four cycles (about 8 to 10% of patients), or five cycles (1.5%) of alemtuzumab. We performed a scenario analysis by varying the proportion of patients who need more than 2 cycles of alemtuzumab during a 5-year period. The results showed that alemtuzumab still was the dominant strategy.

We assumed fixed discontinuation rate across all treatment alternatives for the first two years. We performed scenario analyses to test different discontinuation rates. The results showed that discontinuation rate did not have a significant impact on the results.

We assumed that the average length of mild or moderate relapses was 45 days, and 90 days for severe relapses. The duration of the relapse might be shorter depending on the response to the treatment with corticosteroids. We conducted a scenario analysis where the average length of moderate and severe relapse were 21 days and 45 days. Although some changes in the results were observed, the conclusion remain the same.

The results of our systematic review showed no significant differences between the therapies for serious adverse events. However, the risk of developing progressive multifocal leukoencephalopathy (PML) associated with natalizumab, even if it is rare, was considered important, and, therefore, included in the model. We assumed that the costs and disutility related to other adverse events would not have a significant impact on the results. It should also be mentioned that recently PML has also been reported in some patients treated with other disease-modifying therapies, such as dimethyl fumurate and fingolimod.

The costs associated with inpatient treatment of PML were estimated based on prices from the Norwegian DRG system (DRG code 421). As the costs of inpatient treatment of PML might be underestimated, we performed a scenario analysis where the costs were 100% increased. As the risk of developing PML is low, the correction factor had no significant impact on the cost-effectiveness results.

We performed the health economic evaluation from a health care perspective. The health care perspective is relevant for prioritisation of interventions within a fixed budget if the aim of the decision maker is to maximize health.

Glatiramer acetate 20 mg was included in the base-case analysis. Based on the results from our systematic review regarding relative rates of annual relapse and relative risk of disability progression, and also the estimated annual drug costs, it is highly probable that glatiramer acetate 40 mg 3 times per week will be as cost-effective as glatiramer acetate 20 mg per day (given that all the other parameters are the same).

Due to the uncertain evidence regarding the potential added value of peroral drug administration and the probable effect of the different methods of administrating the medication on patients' utility, we did not include these parameters in the model.

The budget impact estimates were based on several factors that can vary such as disease prevalence and incidence, current clinical practice, drug and healthcare. The market share forecasts for the next five years in our analysis were estimated based on the results of our cost-effectiveness analysis and the drugs' adverse events. We also took under consideration the current practice where there is a trend in favour of oral medicines.

Consistency

Consistency of the systematic review with other publications

Our results are consistent with the results of the Canadian HTA report on drug therapies for RRMS (27), although we included more up to date evidence, and also evidence on more MS treatments. Our results are also consistent with a recently published Cochrane systematic review (93).

Consistency of the economic evaluation with other studies

While several cost-effectiveness studies have examined disease-modifying therapies for RRMS patients, to date, only the Canadian report (27) has compared almost all drugs in one analysis, as we have done in this report. However, it should be mentioned that peg-interferon beta-1a was not included in the Canadian report, and the pricing of alemtuzumab and teriflunomide was not available in Canada at the time the analyses were conducted. Therefore, they were not included in the Canadian base-case analysis.

The Canadian base-case analysis showed that glatiramer acetate was the most costeffective treatment unless willingness to pay exceeded CAD 118,242 per QALY. Between CAD 118,242- CAD 425,655, interferon beta-1b was the most cost-effective treatment, between CAD 425,655- CAD 872,972 it was dimethyl fumarate, and above CAD 872,972, it was natalizumab. It is difficult to compare our results to the Canadian results, as we included more treatment strategies, and used different input data (efficacy, costs and quality of life data).

Conclusion and implications on practice

All examined treatments were more effective than placebo against annual relapse. The strongest effect was seen for alemtuzumab 12 mg. Fingolimod oral 0.5 mg and dimethyl fumarate 240 mg two times a day were also associated with a reduction in annualised relapse rate. For disability progression, direct evidence of high quality indicated that dimethyl fumarate 240 mg twice daily and fingolimod oral 0.5 mg were more effective than placebo. For withdrawal due to adverse events, the lower quality of the available evidence provides unclear conclusion. Results indicate that some treatments are associated with more withdrawal due to adverse events than placebo, such as interferon beta-1a 44 mcg, and all regimens of peg-interferon beta-1a mcg. These results should be considered bearing in mind that some of them are first line treatments while others are used as second line treatments, and may not be relevant to whole type of MS patients.

Our health economic analysis indicated that alemtuzumab dominated all other disease-modifying therapies, as it was more effective and less costly than the other treatment alternatives.

A scenario analysis that excluded alemtuzumab (the dominant strategy) showed that three treatment alternatives (interferon beta-1b (Extavia), peg-interferon beta-1a and natalizumab) could be cost-effective depending on the willingness-to-pay (WTP) threshold. Interferon beta-1b was likely to be the cost-effective choice for a WTP per QALY below NOK 1,658,000. Peg-interferon was the cost-effective option for a WTP from NOK 1,658,450 to NOK 10,619,960, and natalizumab was the cost-effective alternative for a WTP above NOK 10,619,960. Assuming a WTP below NOK 1,000,000 per QALY, interferon beta-1b (Extavia) was approximately 40% likely to be the most cost-effective treatment, followed by peg-interferon beta-1a (approximately 30% likely).

Our budget impact analysis showed that there is a substantial potential for cost saving.

Need for further research

The length of included RCTs is relatively short with a maximum of 3.5 years. We need longer studies to be able to conclude on the longh term efficacy and safety of MS medicines.

Study designs of published studies do not allow to investigate separately first and second line treatments, or to conclude on the sequential use of first and second line treatments. It is difficult to conclude which medicine is most effective when interested only in first or second line treatments. To address this, future studies should use appropriate study design that fits the type of the investigated treatment. For example, first line treatments should be examined as first-line (i.e. in treatment naïve patients), and second line treatments should be investigated as second-line treatments (that is in treatment experienced patients).

There is some degree of uncertainty regarding the health economic model input parameters. More research on efficacy and epidemiologic input parameters would have the greatest impact on reducing decision uncertainty.

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Appendix

Appendix 1: Literature search strategy

Search strategy - Drugs for multiple sclerosis

Databases:	Ovid MEDLINE(R), Embase (Ovid). Cochrane Library: Cochrane Da- tabase of Systematic Reviews, Other Reviews (DARE), Cochrane Cen- tral Register of Controlled Trials (Central), Health Technology As- sessments (HTA), Economic Evaluations (NHS EED). Centre for Reviews and Dissemination: DARE, HTA, NHS EED. Web of Science, PubMed, SweMed+, SBU, Google scholar, PROS-
Date:	PERO. 2015.02.26.
	1.09 updated search for RCT
Study designs	
	izes specificity)" and text words: ((systematic* or literature) adj2 (re-
	view* or overview*)) in title or abstract. Search fliter Ovids "therapy
	(maximizes specificity)" and search filters for RCT's from Cochrane
	Handbook, chapter 6.4.11.1/2.
Limits:	2013-2015 - Randomized controlled trials
Results:	1613 records (277 SR + 729 RCT +607 Econ. Eval.) without duplicates
277 SF	
729 R0	CT (644 + 85 in update search)
607 Eo	conomic evaluations
Searched by:	Ingrid Harboe, research librarian

Search strategies:

Scarch strat	
Databases:	Embase 1974 to 2015 February 25,
	Ovid MEDLINE(R) In-Process & Other Non-Indexed Citations,
	Ovid MEDLINE(R) Daily, Ovid MEDLINE(R) and Ovid
	OLDMEDLINE(R) 1946 to Present
Date:	2015.02.25
Codes:	Embase: oemezd
MEDL	INE: pmoz
SR	
Results:	816 RCT + 69 (update search)

Searches

Results

1	Multiple sclerosis/ or Multiple sclerosis, chronic progressive/ or Multiple sclerosis, relapsing-remitting/ or Neuromyelitis Optica/ use pmoz [Medline]	130140
2	Multiple sclerosis/ use oemezd [Embase]	84701
3	((multiple or disseminated) adj sclerosis).tw.	124063
4	(sclerosis multiplex or Neuromyelitis Optica).tw.	5340
5	((progressive or relapsing or remitting or aggressive or inflammatory or ac- tive) adj MS).tw.	9306
6	(SPMS or PPMS or RRMS).tw.	7859
7	MS.ti.	48528
8	or/1-7	195757
9	Fumaric acid dimethyl ester/ use oemezd	1068
10	(dimethyl fumarate* or dimethylfumarate*).tw.	1054
11	Teriflunomide/ use oemezd	1128
12	teriflunomide.tw.	502
13	Interferon-beta/ use pmoz	7464
14	Beta interferon/ use oemezd	17923
15	(interferon adj1 beta*).tw.	16726
16	Glatiramer/ use oemezd	5518
17	(glatirameracetat* or glatiramer acetat*).tw.	3213
18	Natalizumab/ use oemezd	5744
19	natalizumab.tw.	3941
20	Fingolimod/ use oemezd	4436
21	fingolimod.tw.	2150
22	Alemtuzumab/ use oemezd	10765
23	alemtuzumab.tw.	5127
24	or/9-23	57825
25	8 and 24	19920
26	limit 25 to "reviews (maximizes specificity)"	229
27	((systematic* or literature) adj2 (review* or overview*)).ti,ab.	347467
28	25 and 27	236
29	or/26,28	352

30 lim	it 29 to yr="1995 -Current"	350
31 exp	animals/	37620453
32 hur	nans/	29132069
33 31 1	not (31 and 32)	8488384
34 25	not 33	19194
35 lim	it 34 to "therapy (maximizes specificity)"	1986
36 ran	domized controlled trial.pt. use pmoz	385465
37 con	trolled clinical trial.pt. use pmoz	88645
38 ran	domized.ti,ab. use pmoz	331972
39 pla	cebo.ab. use pmoz	158299
40 clin	ical trials as topic.sh. use pmoz	170938
41 ran	domly.ab. use pmoz	224453
42 tria	l.ti. use pmoz	133387
43 or/	36-42	940316
44 34	and 43	1211
45 ran	domized controlled trial/ use oemezd	363421
46 cro	ssover-procedure/ use oemezd	41657
47 dou	ıble-blind procedure/ use oemezd	120547
48 sing	gle-blind procedure/ use oemezd	19566
49 ran	domized.ab. use oemezd	417485
50 pla	cebo.ab. use oemezd	206226
51 ran	domly.ab. use oemezd	282429
52 tria	l.ti. use oemezd	176165
53 or/	45-52	974635
54 34	and 53	2056
55 35	or 44 or 54	3363
56 lim	it 55 to yr="2013 -Current"	816
	5d or eq-5d or euroqol or euro qol or euroqol-eq-5d or eq-5d-euroqol or 5d-3L or eq-5d-5L).mp.	12866
58 (qu	ality adjusted life or quality-adjust-life).mp.	26318

59	(qaly* or qald* or qale* or qtime* or qali*).mp.	15888
60	57 or 58 or 59	40089
61	25 and 60	249
62	limit 61 to yr="2013 -Current"	69
63	remove duplicates from 56	692
64	"Cost Benefit Analysis"/	128162
65	"Cost Effectiveness Analysis"/	165316
66	"Cost Minimization Analysis"/	44712
67	"Cost Utility Analysis"/	67265
68	(cost* adj2 (analys* or benefit* or effective* or minim* or utilit*)).tw.	246501
69	cba.tw.	19501
70	cea.tw.	41311
71	cua.tw.	1829
72	Economic Evaluation/	71524
73	Health economics/	34220
74	(health economic? or economic evaluation?).tw.	24738
75	Pharmacoeconomics/	8587
76	((pharmacoeconomic? or pharmac*) adj economic?).tw.	863
77	(15D or HRQoL or health-related quality of life instrument).mp.	23802
78	or/60,64-77	541256
79	25 and 78	799
80	Cost-Benefit Analysis/	128162
81	(cost* adj2 (analys* or benefit* or effective* or minim* or utilit*)).tw.	246501
82	cba.tw.	19501
83	cea.tw.	41311
84	cua.tw.	1829
85	Economics, Medical/	42830
86	(health economic? or economic evaluation?).tw.	24738
87	Economics, Pharmaceutical/	8587
88	(pharmac* adj economic?).tw.	863

	89	pharmacoeconomic?.tw.		8935
	90	Technology Assessment, Biomedical/		19671
	91	technology assessment?.tw.		8787
	92	(15D or HRQoL or health-related quality of life instrument).mj	p.	23802
	93	or/60,80-92		489726
	94	25 and 93		736
	95	79 or 94		840
	96	remove duplicates from 95		698
	97	96 not 63		654
	98	97 use oemezd		606
	99	97 use pmoz		48
10	00	limit 56 to yr="2015 -Current"		69
10	01	remove duplicates from 100		62
	02	101 use oemezd		7
	03	101 use pmoz		55
Date	e Run ilts: 2 20 37 41	e: Cochrane Library : 2015.02.26. 24 Cochrane Reviews (Reviews and Protocols), 9 Other Reviews, 7 Technology Assessments Economic Evaluations Clinical trials + 29 (update search)		
#15	Seat MeS MeS MeS ((mu "n #1 0 (dim terif MeS (inte (glat nata fingo alem #7 0		m only 426 ex) or	

#18	 #16 in Cochrane Reviews (Reviews and Protocols), Other Reviews, Technology Assessments and Economic Evaluations #16 Publication Year from 2013 to 2015, in Trials #16 Publication Year from 2015 to 2015, in Trials 	122 181 29
	abase: Centre for Reviews and Dissemination (CRD)	
Date Rest	• /	
Lin	46 NHS EED (Econ. eval.) e Search	Hits
1	MeSH DESCRIPTOR Multiple Sclerosis	201
2	MeSH DESCRIPTOR Multiple Sclerosis, Chronic Progressive	12
3	MeSH DESCRIPTOR Multiple Sclerosis, Relapsing-Remitting	60
4	MeSH DESCRIPTOR Neuromyelitis Optica	1
5	((multiple sclerosis OR disseminated sclerosis OR sclerosis multi- plex OR "neuromyelitis optica"))	408
6	((MS OR SPMS OR PPMS OR RRMS))	808
7	#1 OR #2 OR #3 OR #4 OR #5 OR #6	1052
8	((dimethyl fumarate* or dimethylfumarate*))	12
9	(teriflunomide*)	8
10	MeSH DESCRIPTOR Interferon-beta	68
11	((interferon next beta*))	94
12	((glatiramer aceta* or glatirameraceta*))	32
13	(natalizumab)	34
14	(fingolimod)	22
15	(alemtuzumab)	34
16	#8 OR #9 OR #10 OR #11 OR #12 OR #13 OR #14 OR #15	178
17	#7 AND #16	129
18	(#17) IN DARE, HTA	83
19	(#17) IN NHSEED	46
-)		1~

Database: PubMed

Date: 2015.02.26 **Results: 10 Reviews** 7 RCT + 11 (update search) Search: SR: ((((multiple sclerosis[MeSH Terms]) OR (("multiple sclerosis" OR "disseminated sclerosis" OR "sclerosis multiplex" OR "neuromyelitis optica")) OR "MS" OR SPMS OR PPMS OR RRMS)))) AND ((((((("dimethyl fumarate"[Title/Abstract] OR dimethylfumarate[Title/Abstract]))) OR teriflunomide[Title/Abstract]) OR (("interferon beta"[Title/Abstract]) OR interferon-beta[Title/Abstract]))) OR (("glatiramer aceta"[Title/Abstract] OR glatirameraceta[Title/Abstract]))) OR natalizumab[Title/Abstract]) OR fingolimod[Title/Abstract]) OR alemtuzumab[Title/Abstract])) AND review AND Pubstatusaheadofprint RCT: (((randomized[Title/Abstract] OR randomly[Title/Abstract]))) AND (((((multiple sclerosis[MeSH Terms]) OR (("multiple sclerosis" OR "disseminated sclerosis" OR "sclerosis multiplex" OR "neuromyelitis optica")) OR "MS" OR SPMS OR PPMS OR RRMS)))) AND ((((((((("dimethyl fumarate"[Title/Abstract] OR dimethylfumarate[Title/Abstract]))) OR teriflunomide[Title/Abstract]) OR (("interferon beta"[Title/Abstract] OR interferon-beta[Title/Abstract]))) OR (("glatiramer aceta"[Title/Abstract] OR glatirameraceta[Title/Abstract]))) OR natalizumab[Title/Abstract]) OR fingolimod[Title/Abstract]) OR alemtuzumab[Title/Abstract])) AND pubstatusaheadofprint)

Web of Science Date: 2015.02.26 Results: 11 clinical 53 reviews	
# 16 66	#15 AND #14 Timespan=2013-2015 Search language=Auto
mately	YEAR PUBLISHED: (2013-2015) Timespan=2013-2015 Search language=Auto
# 14 730	#2 AND #1 Refined by: Databases: (WOS) AND Databases: (WOS) AND DOCUMENT TYPES: (CLINICAL TRIAL) <i>Timespan=1995-2015</i> <i>Search language=Auto</i>
# 13 Approxi- mately 14,598	#2 AND #1 Refined by: Databases: (WOS) AND Databases: (WOS) <i>Timespan=1995-2015</i> <i>Search language=Auto</i>
# 12 11	#9 AND #4 Refined by: Databases: (WOS) AND DOCUMENT TYPES: (CLINICAL TRIAL) <i>Timespan=2013-2015</i> <i>Search language=Auto</i>

# 11	50	#9 AND #4 Refined by: Databases: (WOS) <i>Timespan=2013-2015</i> <i>Search language=Auto</i>
# 10	50	#9 AND #4 Timespan=2013-2015 Search language=Auto
#9	Approxi- mately 113,246	TOPIC: (("randomized controlled trial" or randomized* or randomly or "controlled clinical trial")) <i>OR</i> TITLE: (("randomized controlled trial" or randomized* or randomly or "controlled clinical trial")) <i>Timespan=2013-2015 Search language=Auto</i>
# 8	53	#5 AND #4 Refined by: Databases: (WOS) AND DOCUMENT TYPES: (REVIEW) <i>Timespan=1995-2015</i> <i>Search language=Auto</i>
#7	68	#5 AND #4 Refined by: Databases: (WOS) <i>Timespan=1995-2015</i> <i>Search language=Auto</i>
# 6	68	#5 AND #4 Timespan=1995-2015 Search language=Auto
# 5	Approxi- mately 181,139	TOPIC: (systematic* review*) <i>OR</i> TITLE: (systematic* review*) <i>Timespan=1995-2015</i> <i>Search language=Auto</i>
#4	Approxi- mately 14,598	#2 AND #1 Refined by: Databases: (WOS) <i>Timespan=1995-2015</i> <i>Search language=Auto</i>
#3	Approxi- mately 15,657	#2 AND #1 Timespan=1995-2015 Search language=Auto
# 2	Approxi- mately 266,458	TOPIC: (("dimethyl fumarate" OR dimethylfumarate OR teri- flunomide OR interferon OR glatirameraceta* OR "glatiramer aceta" OR natalizumab OR alemtuzumab)) <i>OR</i> TITLE: (("di- methyl fumarate" OR dimethylfumarate OR teriflunomide OR interferon OR glatirameraceta* OR "glatiramer aceta" OR na- talizumab OR alemtuzumab)) <i>Timespan=1995-2015</i> <i>Search language=Auto</i>
#1	Approxi- mately 113,294	TOPIC: (("multiple sclerosis" OR "disseminated sclerosis" OR "sclerosis multiplex" OR "neuromyelitis optica")) <i>OR</i> TI- TLE: (("multiple sclerosis" OR "disseminated sclerosis" OR "sclerosis multiplex" OR "neuromyelitis optica")) <i>Timespan=1995-2015</i> <i>Search language=Auto</i>

Database: PROSPERO

Date: 2015.02.20. Results: 1 Search: multiple sclerosis

Database: SweMed+

Date: 2015.02.20. Results: 8 Search: Multiple sclerosis AND ("dimethyl fumarate" OR dimethylfumarate OR teriflunomide OR interferon OR glatirameraceta* OR "glatiramer aceta*" OR natalizumab OR alemtuzumab)

Webpage: SBU

Date: 2015.02.20. Results: 0 Search: Multipel sckleros

Webpage: Google scholar

Date: 2015.02.20. Results: 2 Search: "Multiple sclerosis" AND name of the intervention drugs AND "technology assessment" AND allintitle "Multiple sclerosis" AND name of the intervention drugs AND "systematic review" AND allintitle

Notes on the following tables:

- Unless otherwise stated, the baseline characteristics described are those of all participants in the study
- Unless otherwise stated, the statistics presented for age and Expanded Disability Status Scale (EDSS) are means (+/-standard deviation)
- The following tables are presented by alphabetic order of the medicine considered as the intervention of interest.
- List of abbreviations used in tables:
 - IV= intravenous;
 - IM= intra muscular
 - SC= subcutaneous;
 - mg = milligram
 - mcg=micrograms
 - q.d.= once daily q.w.= once weekly
 - t.i.w.= three times weekly

Alemtuzumab

CAMMS223-study 2008,	CAMMS223 Tria	l Investigators	(28), included
(incl.) in Khai et al. (27)		_	

	-	
RCT identification	NCT00050778	
Study setting	Rater-blinded, and US	randomized controlled trial in 49 centres in Europe
Participants	Eligibility crite	eria: Diagnosis of RRMS (McDonald criteria) with an
_		toms no more than 36 months before the time of
	screening, EDS	SS = 0 to 3.0; had one or more enhancing lesions on
		relapses during the previous 2 years.
		criteria: Previous disease-modifying treatment; pres-
		antithyrotropin-receptor antibodies.
		acteristics: Age 32+/-8; 64% female; EDSS 2,0+/-0.8
Intervention group	Annual alemtu	
		o 12 mg IV q.d., 5 consecutive days at 1st month, 3 con-
		t months 12 and 24 (n = 113)
~ •		p 24 mg IV q.d. (n = 110)
Comparison group		a-1a 44 mcg SC t.i.w. (n = 111)
Outcome	0 1	<u>pints:</u> Sustained accumulation of disability and rate of
	relapse.	herinte. Drementies of matients with malance from MO
		<i>lpoints</i> : Proportion of patients with relapse-free MS,
	different MRI o	
		ed for endpoints: <i>Relapses</i> : New or worsening symp- objective change in neurologic examination attributa-
		a lasted 48 hours, that were present at normal body
		and that were preceded by at least 30 days of clinical
	stability.	ind that were preceded by at least 30 days of chinical
		umulation of disability: An increase of at least 1.5
		ents with baseline score of 0, and at least 1.0 point for
		a baseline score of 1.0 or more; all scores were con-
		uring a 6-month period.
Follow-up	3 years	
Treatment history		ve (based on inclusion criteria)
Comments		2005, alemtuzumab therapy was suspended after im-
	mune thrombo	cytopenic purpura developed in three patients, one of
	whom died. Tre	eatment with interferon beta-1a continued throughout
	the study.	
Critical appraisal		
Randomization		Adequate
Allocation concealment	nt	Insufficient reporting
Double-blinding		No (rater-blinded)
Baseline characteristic	c similarity	Yes
Outcome measures		Adequate
Withdrawals		25%
· · · · · · · · · · · · · · · · · · ·	ITT Analysis Yes	
Funding	ng Manufacturer	

CARE (Comparison of Alemtuzumab and Rebif Effi cacy in Multiple Sclerosis) MS I- study 2012, Cohen et al. (29), in Khai et al. (27)

countries including Europe, CanParticipantsEligibility criteria: Age = 18 ye (McDonald criteria) with disease 3.0; had cranial abnormalities of relapses during the previous 2 ye Key exclusion criteria: Progress ease therapy (apart from corticos sive; investigational or monoclo nificant autoimmunity other that	ars to 50 years, diagnosis of RRMS e duration up to 5 years, EDSS = 0 to on MRI attributable to MS; with ≥ 2 ears. ive disease course, previous MS dis- steroids), previous immunosuppres- nal antibody therapy, clinically sig- n MS. +/-8; 65% female; EDSS 2.0+/-0.8 consecutive days at month 0, 3 con-		
countries including Europe, CanParticipantsEligibility criteria: Age = 18 ye (McDonald criteria) with disease 3.0; had cranial abnormalities of relapses during the previous 2 ye Key exclusion criteria: Progress ease therapy (apart from corticos sive; investigational or monoclo nificant autoimmunity other that	ada, and US. ars to 50 years, diagnosis of RRMS e duration up to 5 years, EDSS = 0 to on MRI attributable to MS; with ≥ 2 ears. ive disease course, previous MS dis- steroids), previous immunosuppres- nal antibody therapy, clinically sig- n MS. +/-8; 65% female; EDSS 2.0+/-0.8 consecutive days at month 0, 3 con-		
ParticipantsEligibility criteria: (McDonald criteria) with disease 3.0; had cranial abnormalities of relapses during the previous 2 you Key exclusion criteria: Progress ease therapy (apart from corticos) sive; investigational or monoclon inficant autoimmunity other that	ars to 50 years, diagnosis of RRMS e duration up to 5 years, EDSS = 0 to on MRI attributable to MS; with ≥ 2 ears. ive disease course, previous MS dis- steroids), previous immunosuppres- nal antibody therapy, clinically sig- n MS. +/-8; 65% female; EDSS 2.0+/-0.8 consecutive days at month 0, 3 con-		
(McDonald criteria) with disease 3.0; had cranial abnormalities of relapses during the previous 2 yo <u>Key exclusion criteria</u> : Progress ease therapy (apart from corticos sive; investigational or monoclo nificant autoimmunity other tha	e duration up to 5 years, EDSS = 0 to on MRI attributable to MS; with ≥ 2 ears. ive disease course, previous MS dis- steroids), previous immunosuppres- nal antibody therapy, clinically sig- n MS. +/-8; 65% female; EDSS 2.0+/-0.8 consecutive days at month 0, 3 con-		
relapses during the previous 2 ye <u>Key exclusion criteria</u> : Progress ease therapy (apart from corticos sive; investigational or monoclo nificant autoimmunity other tha	ears. ive disease course, previous MS dis- steroids), previous immunosuppres- nal antibody therapy, clinically sig- n MS. +/-8; 65% female; EDSS 2.0+/-0.8 consecutive days at month 0, 3 con-		
<u>Key exclusion criteria:</u> Progress ease therapy (apart from corticos sive; investigational or monoclo nificant autoimmunity other tha	ive disease course, previous MS dis- steroids), previous immunosuppres- nal antibody therapy, clinically sig- n MS. +/-8; 65% female; EDSS 2.0+/-0.8 consecutive days at month 0, 3 con-		
ease therapy (apart from corticos sive; investigational or monoclo nificant autoimmunity other tha	steroids), previous immunosuppres- nal antibody therapy, clinically sig- n MS. +/-8; 65% female; EDSS 2.0+/-0.8 consecutive days at month 0, 3 con-		
ease therapy (apart from corticos sive; investigational or monoclo nificant autoimmunity other tha	steroids), previous immunosuppres- nal antibody therapy, clinically sig- n MS. +/-8; 65% female; EDSS 2.0+/-0.8 consecutive days at month 0, 3 con-		
nificant autoimmunity other tha	n MS. +/-8; 65% female; EDSS 2.0+/-0.8 consecutive days at month 0, 3 con-		
	+/-8; 65% female; EDSS 2.0+/-0.8 consecutive days at month 0, 3 con-		
Baseline characteristics: Age 33	consecutive days at month 0, 3 con-		
	consecutive days at month 0, 3 con-		
Intervention group Alemtuzumab 12 mg IV q.d., 5 d	Alemtuzumab 12 mg IV q.d., 5 consecutive days at month 0, 3 con-		
secutive days at month 12 (n = 3			
Comparison group Interferon beta-1a 44 mcg SC t.i			
	e and time to 6 months sustained ac-		
cumulation of disability.			
	ion of patients with relapse-free,		
change in EDSS, change in MSF	C, different MRI outcomes.		
Definitions used for endpoints:	<i>Relapses:</i> New or worsening neuro-		
	IS, lasting at least 48 hours, with py-		
	clinical stability, with an objective		
change on neurological examina			
	<i>bility</i> : An increase from baseline of		
	5 points if baseline EDSS score was		
o) confirmed over 6 months.	-		
Follow-up2 years			
Treatment history Treatment-naive (based on inclu	ision criteria).		
Critical appraisal			
RandomizationAdequate			
Allocation concealment Adequate			
Double-blinding No (rater-blinde	ed)		
Baseline characteristic similarity Yes			
Outcome measures Adequate			
Withdrawals 9%			
ITT Analysis Yes			
FundingManufacturer			

CARE (Comparison of Alemtuzu mab and Rebif Effi cacy in Multiple Sclerosis)-MS II study 2012, Coles et al. (30), in Khai et al. (27)

	NOTeer () (er		
RCT identification	NCT00548405		
Study setting		randomized controlled trial. 194 academic medical	
		inical practices in 23 countries including Europe,	
	Canada, and U		
Participants		ria: Age = 18 years to 50 years, diagnosis of RRMS	
		teria) with disease duration up to 5 years, $EDSS = 0$	
		nial and spinal MRI lesions; with ≥ 2 relapses dur-	
		s 2 years and at least one in the previous year.	
		<u>criteria</u> : Progressive forms of MS, previous cyto-	
	toxic drug use or investigational therapy, treatment within the previous 6 months with natalizumab, methotrexate, azathioprine		
		e, and a history of clinically significant autoimmun-	
	ity other than M		
	•	<i>cteristics:</i> Age: 35 +/-8, 67 female, EDSS: 2.7 +/-	
	1.2	<u>elevisites.</u> Age: 55 +/ 6, 6/ lemaic, 1065. 2./ +/	
Intervention group		12 mg IV q.d., 5 consecutive days at month 0, 3	
		ys at month 12 (n=436)	
		24 mg IV q.d., 5 consecutive days at month 0, 3	
		ys at month 12 (n=173)	
Comparison group	Interferon beta	1a 44 mcg SC t.i.w. (n=231)	
Outcome	Primary endpo	<i><u>pints:</u></i> Relapse rate and time to 6 months sustained	
	accumulation of		
		<i>dpoints</i> : Proportion of patients with relapse-free,	
	change in EDSS, change in MSFC, different MRI outcomes.		
	Definitions used for endpoints: Relapses: New or worsening neu-		
	rologic symptoms attributable to MS, lasting at least 48 hours,		
	without pyrexia, after at least 30 days of clinical stability, with an		
	objective change on neurological examination.		
	Sustained accumulation of disability: An increase from baseline of		
	at least one EDSS point (or \geq 1.5 points if baseline EDSS score was		
	o) confirmed over 6 months.		
Follow-up	2 years		
Treatment history	Treatment-experienced (based on inclusion criteria).		
Comments	The 24 mg per day group was discontinued to aid recruitment, but		
	data are included for safety assessments		
Critical appraisal			
Randomization	+	Adequate	
Allocation concealmen	ll .	Adequate No (rater blinded)	
Baseline characteristic	similarity	Yes	
Outcome measures	siiiiafity	Adequate	
Withdrawals		15%	
ITT Analysis		Yes	
Funding		Manufacturer	
runuing M			

Dimetyl fumarate

DEFINE (Determination of the Efficacy and Safety of Oral Fumarate in Relapsing–Remitting MS) study, Gold 2012 (33), in Khai et al. (27)

RCT identification	NCT00420212	
Study setting		puble-blind, placebo controlled trial. 198 sites in 28
		ing Europe, Canada, and US
Participants		<u>ria:</u> Age = 18 years to 55 years, diagnosis of RRMS
		eria), EDSS = 0 to 5.0; \geq 1 clinically documented re-
	lapse within 12	months before randomization, or \geq 1 gadolinium-en-
	hancing lesion w	vithin 6 weeks before randomization
	Key exclusion criteria: Progressive forms of MS, another major dis-	
	ease that would	preclude participation in the clinical trial, abnormal
	results on the pre-	e-specified laboratory tests, or recent exposure to con-
	traindicated med	
	Baseline chara	cteristics: Age: 38+/-9 years; 74% female; EDSS
	2,4+/-1,2	
Intervention group		ate 240 mg oral twice daily (480 mg/day) (n = 410)
0 1		ate 240 mg oral 3 times daily (720 mg/day) $(n = 416)$
Comparison group	Placebo $(n = 408)$	
Outcome		<i>nt</i> : Patients' proportion who had a relapse by 2 years
		<u>oints:</u> Different MRI outcomes at 2 years, annualized
		e to progression disability.
	-	
	<u>Definitions used for endpoints:</u> <u><i>Relapses</i></u> : New or recurrent neuro- logic symptoms, not associated with fever or infection, that lasted at	
	least 24 hours and that were accompanied by new objective neuro-	
	logic findings according to neurologist's evaluation.	
	<u>Disability progression:</u> At least a 1.0-point increase on the EDSS in	
	patients with a baseline score of 1.0 or higher or at least a 1.5-point	
	increase in patients with a baseline score of 0, with the increased score	
	sustained for at least 12 weeks.	
Follow-up	2 years	
Treatment history	Mixed (based on baseline characteristics)	
Comments	Patients could switch to an approved alternative MS therapy if they	
Comments		
	had completed 48 weeks of blinded treatment, and had at least 1 con-	
	firmed relapse after 24 weeks, or at any time if they had experienced disability progression sustained for 12 weeks.	
Critical appraisal	uisability progre	ssion sustained for 12 weeks.
Randomization		Adequate
Allocation concealme	nt	Adequate
Double-blinding	/110	Yes
Baseline characterist	ic similarity	Yes
Outcome measures	it similarity	Adequate
Withdrawals		1
		23% Voc
ITT Analysis		Yes (Discord)
Funding		Manufacturer (Biogen)

CONFIRM (Comparator and an Oral Fumarate in Relapsing–Remitting Multiple Sclerosis) study 2012, Fox et al., (34), in Khai et a. *(27)*

RCT identification	NCT00451451	
Study setting	Rater-blinded,	randomized controlled trial. in 200 research sites in
	28 countries in	cluding Europe and North America
Participants	Eligibility crite	eria: RRMS (McDonald criteria), age 18 to 55 years,
	EDSS o to 5 an	nd at least one clinically documented relapse in the
		onths or at least one gadolinium-enhancing lesion o
		pre randomization.
		<i>criteria:</i> Progressive forms of multiple sclerosis,11
		v significant illness, prespecified laboratory abnor-
		prior exposure to glatiramer acetate or contraindi-
	cated medicati	
	<u>Baseline charc</u>	<i>acteristics:</i> Age: 37 +/-9, 70% female, EDSS score:
	2.6 +/-1.2	
Intervention group		rate 240 mg b.i.d, (n=359)
		rate 240 mg three times daily (n=345), subcutane-
		tions of 20 mg of glatiramer acetate for 96 weeks
	(n=350)	
Comparison group	Placebo (n=36)	
Outcome	Primary endpoint: Annualized relapse rate at 2 years.	
		<i>lpoints</i> : Different MRI outcomes at 2 years, disabil-
	ity progression	
	Tertiary endpoints: Relative benefits and risks of BG-12 or glati-	
	ramer acetate versus placebo and the number of gadolinium-en-	
	hancing lesions at 2 years.	
	Definitions use	ed for endpoints: <i>Relapses:</i> New or recurrent neuro-
		s not associated with fever or infection, lasting at
		accompanied by new objective neurologic findings,
	and separated from the onset of other confirmed relapses by at least	
	30 days.	
	Disability progression: An increase in the EDSS score of at least 1.0	
	point in patients with a baseline score of 1.0 or more or an increase	
	of at least 1.5 points in patients with a baseline score of 0, confirmed	
	at least 12 weeks later.	
Follow-up	2 years	
Treatment history	Mixed (based on reported baseline characteristics)	
Critical appraisal		
Randomization		Adequate
Allocation concealeme	nt	Adequate
Double-blinding		No
Baseline characteristic	similarity	Yes
Outcome measures		Adequate
Withdrawals		21%
ITT Analysis		Yes
Funding		Manufacturer (Biogen Idec)

Fingolimod

FREEDOMS (FTY720 Research Evaluating Effects of Daily Oral Therapy in Multiple Sclerosis) study, Kappos 2010 (37), in Khai et al. (27)

RCT identification	NCT00289978		
Study setting	Double-blind,	randomized, placebo-controlled trial multi-centre in	
	Australia, Can	ada, Europe, and South Africa (138 centers in 22	
	countries)		
Participants		<u>eria:</u> Age = 18 years to 55 years, diagnosis of RRMS	
		teria), EDSS = 0 to 5.5 ; ≥ 1 relapse in the previous	
		pses in the previous 2 years.	
		criteria: Relapse or corticosteroid treatment within	
	30 days before randomization, active infection, macular edema, di-		
	abetes mellitus, immune suppression (drug- or disease-induced),		
	or clinically significant systemic disease.		
		<i>cteristics:</i> Age 37+/-9; 70% female; EDSS 2,4+/-1,4	
Intervention group		al 0.5 mg q.d. (n = 425)	
		al 1,25 mg q.d. (n = 429)	
Comparison group	Placebo $(n = 41)$		
Outcome		<u>pint:</u> Annualized relapse rate.	
		<i>dpoints:</i> Disability progression, time to a first re- ange, MSFC change, different MRI outcomes.	
	lapse, EDSS cli	ange, MSFC change, unterent MKI outcomes.	
	Definitions use	ed for endpoints: <u>Relapses:</u> A confirmed relapse con-	
		oms that must have been accompanied by an in-	
	crease of at least half a point in the EDSS score, of 1 point in each		
		inctional system scores, or of 2 points in one EDSS	
		em score (excluding scores for the bowel-bladder or	
	cerebral function		
		pression: An increase of 1 point in the EDSS score (or	
	half a point if the baseline EDSS score was equal to 5.5), confirmed		
	after 3 months, with an absence of relapse at the time of assessment		
	and with all EDSS scores measured during that time meeting the		
	criteria for disability progression.		
Follow-up	2 years		
Treatment history	Mixed (based on reported baseline characteristics)		
Critical appraisal			
Randomization		Adequate	
Allocation concealment	nt	Adequate	
Double-blinding		Yes	
Baseline characteristic	e similarity	Yes	
Outcome measures		Adequate	
Withdrawals		19%	
ITT Analysis		Yes	
Funding		Manufacturer	

TRANSFORMS (Trial Assessing Injectable Interferon versus FTY720 Oral in Relapsing–Remitting Multiple Sclerosis) study; Cohen et al. 2010, (38), in Khai et al. (27)

2010, (38), III Khai et a				
RCT identification	NCT00340834			
Study setting		randomized controlled trial. 172 centres in 18 coun-		
		Canada, Australia, Europe, and US.		
Participants		eria: Age = 18 years to 55 years; diagnosis of RRMS		
		teria), EDSS = 0 to 5.5; had \geq 1 relapse during the		
		or \geq 2 relapses during the previous 2 years.		
		<u>criteria:</u> Documented relapse or corticosteroid treat-		
		ment within 30 days before randomization; active infection, macu-		
		nunosuppression, and clinically significant coexist-		
	ing systemic di			
	<u>Baseline chara</u>	<u>ucteristics:</u> Age: 36+/-9; 67% female; EDSS: 2.2 +/-		
	1.3			
Intervention group		al 0.5 mg q.d. (n=431)		
		al 1.25 mg q.d. (n=426)		
Comparison group		n-1a 30 mcg IM q.w. (n=435)		
Outcome		<u>pint:</u> Annualized relapse rate.		
		<i>points</i> : Number of new or enlarged T2-hyperintense		
	lesions, time to	confirmed disability progression		
	Definitions used for endpoints: <u>Relapses</u> : New, worsening, or re-			
	current neurologic symptoms that occurred at least 30 days after			
	the onset of preceding relapse, that lasted at least 24 hours without			
	fever or infection.			
	<u>Disability progression:</u> A one-point increase in the EDSS score (or			
	a half-point increase for patients with a baseline score \geq 5.5) that			
	was confirmed 3 months later in the absence of relapse.			
Follow-up	1 year			
Treatment history	Mixed (based on reported baseline characteristics)			
Critical appraisal				
Randomization		Adequate		
Allocation concealment	nt	Adequate		
Double-blinding		Yes		
Baseline characteristi	c similarity	Yes		
Outcome measures		Adequate		
Withdrawals		11%		
ITT Analysis		Yes		
Funding		Manufacturer		

Saida et al. 2012 (39), included in Khai et al. (27)

RCT identification	NCT00537082	2	
Study setting	Double-blind,	randomized controlled trial. Multicentre in Japan	
Participants	<u>Eligibility criteria</u> : Age = 18 years to 60 years, diagnosis of RRMS (McDonald criteria), EDSS = 0 to 6.0; had \geq 1 relapse in the previ- ous year or \geq 2 relapses in the previous 2 years; \geq 1 gadolinium- enhancing lesion within 30 days before study commencement. <u>Key exclusion criteria</u> : Primary-progressive MS; relapse or cortico- steroid treatment within 30 days before randomization; malig- nancy, macular edema, diabetes mellitus, active infection, immu- nosuppression, or significant systemic disease; received cladribine, cyclophosphamide, mitoxantrone, or other immunosuppressive or immunoglobulin medication in the six months before randomiza- tion, or had plasmapheresis immunoadsorption or IFN beta ther- apy in the three months before randomization. <u>Baseline characteristics</u> : Age: 35 +/-9; 69% female; EDSS: 2.1 +/- 1.8		
Intervention group		al 0.5 mg q.d. (n=57)	
	Fingolimod ora	al 1.25 mg q.d. (n=57)	
Comparison group		Placebo (n=57)	
Outcome	<u>Primary endpoint:</u> Percentage of patients free from gadolinium enhanced lesions at 3 and 6 months. <u>Secondary endpoints:</u> Percentage of patients free from relapse over 6 months, annualized relapse rate, and other MRI outcomes. Definitions not reported		
Follow-up	6 months		
Treatment history	Unclear (inadequate information to characterise)		
Critical appraisal			
Randomization		Insufficient reporting	
Allocation concealment	nt	Not reporting	
Double-blinding		Yes	
Baseline characteristic	c similarity	Yes	
Outcome measures		Adequate	
Withdrawals		14%	
ITT Analysis		No	
Funding		Manufacturer	

FREEDOMS II- study (41), not included in Khai et al.(27)

RCT identification	NCT00355134		
Study setting		lomised controlled study. In 117 academic and	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		attres in 8 countries, most patients from USA	
Participants	~ ~	diagnosed with relapsing-remitting multiple	
•		to the 2005 revised McDonald criteria, aged	
	18–55 years, one of	r more confirmed relapses during the preceding	
	year (or two or mor	e confirmed relapses during the previous 2	
	years), EDSS score	of 0-5.5, and had no relapse or steroid treat-	
		$\beta$ s before randomisation. interferon $\beta$ or glati-	
		apy was stopped at least 3 months before ran-	
	domisation and natalizumab treatment at least 6 months before		
	randomisation.		
		<u>ria:</u> clinically significant systemic disease or im-	
	· · · · · · · · · · · · · · · · ·	, active infection or macular oedema, diabetes	
		ry of malignancy, and patients with specific car-	
	diac, pulmonary, or	istics in placebo group: Age: 40+/-8; 81% fe-	
	male; EDSS: 2.4 +/		
Intervention group	Fingolimod 0.5 mg	-	
intervention group	Fingolimod 1.25 mg		
		dose stopped due to absence of clear added ben-	
		safety events risk (infections,macular oedema).	
	e	hed to the 0.5 mg dose in a blinded manner	
Comparison group	Placebo (n=355)		
Outcome		<u>::</u> Annualised relapse rates	
	<u>Secondary endpoints:</u> Percent brain-volume change , the time to		
	first relapse and proportion of relapsefree patients; time to disabil-		
		nfirmed at6 months, as measured by EDSS;	
		ne to the end of study on the MSFC score; and	
	effect on MRI.	r and points: Rolance: confirmed when accom	
	<u>Definitions used for endpoints:</u> <u><i>Relapse:</i></u> confirmed when accompanied by an increase of at least half a step $(0 \cdot 5)$ on the EDSS,		
	an increase of 1 point on two different functional systems of the		
	EDSS, or 2 points on one of the functional systems (excluding bowel, bladder, or cerebral functional systems).		
	<i><u>Disability progression:</u></i> 1 point EDSS change [0· 5 point if base-		
	line EDSS was $>5$ · 0]) confirmed at 3 months for up to 24		
	months.		
Follow-up	2 years		
Treatment history	Unclear (inadequat	e information to characterise)	
Risk of bias			
Random sequence gen		Adequate	
Allocation concealmen		Adequate	
Blinding of participant Blinding of outcome as		Adequate	
Incomplete outcome d		Adequate Intention-to-treat analysis	
	ala	Withdrawals: 28%	
Selective reporting		None detected	
Other sources of bias		Funding: Manufacturer	
Other sources of blas		i ununig, manufacturer	

#### Glatiramer acetate

#### Johnson et al., 1995 (42), included in Khai et al. (27)

RCT identification		
	Not reported	
Study setting	the US	randomized, placebo-controlled trial. 11 centres in
Participants	<u>Eligibility criteria:</u> RRMS (Poser-criteria), age 18 to 45 years, EDSS = 0 to 5.0; had $\geq$ 2 clinically documented relapses in the 2 years before entry; onset of the first relapse at least 1 year before randomization; and a period of neurologic stability and freedom from corticosteroid therapy of at least 30 days prior to entry. <u>Key exclusion criteria:</u> Received Glatiramer acetate 1 or previous immunosuppressive therapy with cytotoxic chemotherapy (azathi- oprine, cyclophosphamide, or cyclosporine) or lymphoid irradia- tion; pregnancy or lactation; insulin-dependent diabetes mellitus, positive HIV or HTL V-I serology, evidence of Lyme disease, or re- quired use of aspirin or chronic nonsteroidal antiinflammatory drugs during the course of the trial. <u>Baseline characteristics</u> : Age: 34+/-6; 73% female; EDSS 2.6 +/- 1.3	
Intervention group	Glatiramer acetate 20 mg SC q.d (n =125)	
Comparison group	Placebo (n=126)	
Outcome	<u><i>Primary endpoints:</i></u> Relapse rate over 24 months, annualized re-	
	<ul> <li>lapse rate, number of relapse over 24 months.</li> <li><u>Secondary endpoints:</u> Proportion of relapse-free patients, median time to first relapse, number of relapse per patient, proportion of patients with a change in disability, EDSS change, proportion of progression-free patients, ambulation index.</li> <li><u>Definitions used for endpoints: <i>Relapses</i>:</u> The appearance or reappearance of one or more neurologic abnormalities persisting for at least 48 hours and immediately proceeded by a relatively stable or improving neurologic state of at least 30 days.</li> <li><u>Disability progression:</u> An increase of at least one full step on the EDSS that persisted of at least 3 months.</li> </ul>	
Follow-up	2 years	
Treatment history	Treatment-naive (based on exclusion criteria, year of study, and clinical expert input).	
Critical appraisal		
Randomization		Insufficient reporting
Allocation concealmer	ıt	Not reporting
Double-blinding		Yes
Baseline characteristic	: similarity	Yes
Outcome measures		Adequate
Withdrawals		14%
ITT Analysis		Yes
Funding		Manufacturer, public

Comi et al., 2001 (43), included in Khai et al. (27)

<b>RCT identification</b>	Not reported		
Study setting	-	randomized controlled study. 29 centres in 6 Euro-	
Study Setting	pean countries		
Participants	1	eria: Age = 18 years to 50 years, with relapse-remit-	
<b>F</b>		diagnosis of MS for at least 1 year, EDSS = 0 to 5.0;	
	•	d relapse in the preceding 2 years, $\geq 1$ enhancing le-	
	sion on screeni		
		criteria: previous use of glatiramer acetate, oral my-	
	elin, lymphoid irradiation, the use of immunosuppressant or cyto-		
		the past 2 years, or the use of azathioprine, cyclo-	
		erons, deoxyspergualine, or chronic corticosteroids	
		vious 6 months.	
		acteristics in placebo group: Age: 34.0+/-8; % fe-	
	male not repor	ted; EDSS: 2,4+/-1.2	
Intervention group	Glatiramer ace	tate 20 mg SC q.d. (n=119)	
Comparison group	Placebo (n=120	o)	
Outcome		<i><u>oint:</u></i> Total number of enhancing lesions.	
	Secondary end	<i>lpoints</i> : Other different MRI outcomes.	
	<u>Tertiary endpo</u>	oints: Relapse rate, percentage of patients with re-	
	lapse-free, steroid courses, relapse-related hospitalizations.		
	Definitions used for endpoints: <i>Relapses</i> : The appearance of one or		
		rological symptoms, or the reappearance of one or	
		more previously experienced ones. An event was counted as a re-	
1			
	lapse only whe	n the patient's symptoms were accompanied by ob-	
		n the patient's symptoms were accompanied by ob- s in the neurological examination corresponding to	
	jective changes		
	jective changes an increase of score of the tw	s in the neurological examination corresponding to at least 0.5 points on the EDSS, or one grade in the o or more functional systems, or two grades in one	
	jective changes an increase of score of the tw functional syste	s in the neurological examination corresponding to at least 0.5 points on the EDSS, or one grade in the o or more functional systems, or two grades in one	
Follow-up	jective changes an increase of score of the tw functional syste 9 months	s in the neurological examination corresponding to at least 0.5 points on the EDSS, or one grade in the ro or more functional systems, or two grades in one em.	
Treatment history	jective changes an increase of score of the tw functional syste 9 months	s in the neurological examination corresponding to at least 0.5 points on the EDSS, or one grade in the o or more functional systems, or two grades in one	
Treatment history Critical appraisal	jective changes an increase of score of the tw functional syste 9 months	s in the neurological examination corresponding to at least 0.5 points on the EDSS, or one grade in the to or more functional systems, or two grades in one em.	
Treatment history Critical appraisal Randomization	jective changes an increase of a score of the tw functional syste 9 months Unclear (inade	s in the neurological examination corresponding to at least 0.5 points on the EDSS, or one grade in the to or more functional systems, or two grades in one em. quate information to characterize) Adequate	
Treatment history Critical appraisal Randomization Allocation concealmer	jective changes an increase of a score of the tw functional syste 9 months Unclear (inade	s in the neurological examination corresponding to at least 0.5 points on the EDSS, or one grade in the ro or more functional systems, or two grades in one em. quate information to characterize) Adequate Adequate	
Treatment history Critical appraisal Randomization Allocation concealmer Double-blinding	jective changes an increase of score of the tw functional syste 9 months Unclear (inade	s in the neurological examination corresponding to at least 0.5 points on the EDSS, or one grade in the ro or more functional systems, or two grades in one em. equate information to characterize) Adequate Adequate Yes	
Treatment history Critical appraisal Randomization Allocation concealmer Double-blinding Baseline characteristic	jective changes an increase of score of the tw functional syste 9 months Unclear (inade	s in the neurological examination corresponding to at least 0.5 points on the EDSS, or one grade in the ro or more functional systems, or two grades in one em. equate information to characterize) Adequate Adequate Yes Yes	
Treatment history Critical appraisal Randomization Allocation concealmer Double-blinding Baseline characteristic Outcome measures	jective changes an increase of score of the tw functional syste 9 months Unclear (inade	s in the neurological examination corresponding to at least 0.5 points on the EDSS, or one grade in the to or more functional systems, or two grades in one em. quate information to characterize) Adequate Adequate Yes Yes Adequate	
Treatment history Critical appraisal Randomization Allocation concealmen Double-blinding Baseline characteristic Outcome measures Withdrawals	jective changes an increase of score of the tw functional syste 9 months Unclear (inade	s in the neurological examination corresponding to at least 0.5 points on the EDSS, or one grade in the ro or more functional systems, or two grades in one em. equate information to characterize) Adequate Adequate Yes Yes Adequate 6%	
Treatment history Critical appraisal Randomization Allocation concealmer Double-blinding Baseline characteristic Outcome measures	jective changes an increase of score of the tw functional syste 9 months Unclear (inade	s in the neurological examination corresponding to at least 0.5 points on the EDSS, or one grade in the to or more functional systems, or two grades in one em. quate information to characterize) Adequate Adequate Yes Yes Adequate	

# REGARD (REbif vs Glatiramer Acetate in Relapsing MS Disease) study 2008, Mikol et al., (44), in Khai et al. (27)

<b>RCT identification</b>	NCT00078338		
Study cotting	Randomized comparative study. Open-label, rater-masked. 81 cen-		
Study setting			
Dearth allow and a		ies (e.g. Canada, South America, and Europe)	
Participants		<i>ia</i> : Adult RRMS patients (McDonald criteria), EDSS	
		1 relapse in the preceding 12 months, and clinically	
		ogically improving during the 4 weeks before ran-	
	domization.		
		<u>riteria:</u> Pregnancy or breastfeeding; treatment with	
		nocorticotropic hormone with the previous 4 weeks;	
	previous treatment with interferon beta, glatiramer acetate, or		
		lymphoid irradiation; plasma exchange within the	
	previous 3 mont	hs; intravenous gamma-globulin use within the pre-	
	vious 6 months;	cytokine or anti-cytokine therapy within the previ-	
		immunosuppressant use within the past 12 months.	
	Baseline charact	<i>teristics:</i> Age: 37+/-10; 71% female; EDSS: 2.3+/-1.3	
Intervention group	Glatiramer aceta	te 20 mg SC q.d. (n=378)	
Comparison group	Interferon beta-	1a 44 mcg SC t.i.w. (n=386)	
Outcome	Primary endpoin	<u>nt:</u> Time to first relapse over 96 weeks.	
	Secondary endp	oints: Mean number T2 active lesions, mean num-	
	ber gadolinium-	enhancing lesions, change in T2 lesion volume.	
		nt: Other MRI outcomes, relapse outcomes, disabil-	
	ity progression.		
	ку рюбловіон.		
	Definitions used for endpoints: <i>Relapses</i> : New or worsening neuro-		
	logical symptoms, without fever, that lasted for 48 hours or more and		
		a change in the Kurtzke Functional Systems Scores.	
		<u>ession</u> : Disability progression at the 6-month fol-	
	low-up visit was confirmed, as follows – if the EDSS score at the		
	baseline was 0, then a change of 1.5 points or more was required; if		
	the EDSS was 0.5 - 4.5 at baseline, then a change of 1.0 point or more		
	was required; and if the EDSS at baseline was 5 points or more, then		
	the change required was 0.5 points or more.		
Follow-up	96 weeks		
Treatment history	Treatment-naive (based on inclusion criteria, year of study, and clin-		
~	ical expert input).		
Critical appraisal			
Randomization		Adequate	
Allocation concealme	nt	Adequate	
Double-blinding		Yes	
Baseline characteristi	c similarity	Yes	
Outcome measures		Adequate	
Withdrawals		18%	
		Yes	
ITT Analysis Funding		Yes	

BECOME (Betaseron vs Copaxone in Multiple Sclerosis with Triple-Dose Gadolinium and 3-Tesla MRI Endpoints) study 2009, Cadavid et al.(45), included in Khai et al. (27)

al. (45), liitluueu ili Kila		
<b>RCT identification</b>	NCT00176592	
Study setting	Rater-blinded,	randomized controlled trial. In one centre in the US.
Participants		<i>ria:</i> Age = 18 years to 55 years; treatment-naïve pa-
	tients with RRM	AS (79%) or CIS (21%) suggestive of MS.
	Exclusion criter	ria: Not reported.
	Baseline chare	acteristics: in interferon beta-1b group: mean
	(range) age 36(18-49); 75% female; EDSS median(range) 2,0 (0-5).	
Intervention group		tate 20 mg SC q.d. (n = 39)
Comparison group	Interferon beta	-1b 250 mcg SC every other day $(n = 36)$
Outcome		ints: Different MRI outcomes at 1 and 2 years. Con-
	firmed relapse	occurrences (annualized relapse rate, percent re-
	lapse-free).	
	_	
	Definitions use	ed for: <u>Relapses:</u> All new or worsening symptoms
		ars and not explained by fever or infection that were
		a blinded examining neurologist using worsening
	scores on SNRS or EDSS. : required for relapse confirmation: 1) in-	
	crease in total EDSS by _0.5 point; 2) increase in the EDSS score	
	for one system _2 points; 3) increase in the score of 2 or more EDSS	
	systems_1 point;	
Follow-up	2 years	
Treatment history	Treatment-naive (based on reported baseline characteristics).	
Critical appraisal		
Randomization		
Allocation concealment	ıt	Not reported
Double-blinding		No (but rater blinded)
<b>Baseline characteristic</b>	similarity	Yes
Outcome measures		Adequate
Withdrawals		15%
ITT Analysis		Yes
Funding		Manufacturer

BEYOND (Betaferon Effi cacy Yielding Outcomes of a New Dose) study 2009, O'Connor et al. (46), included in Khai et al. (27)

RCT identification	NCT00099502	
Study setting		d, randomized controlled trial in 198 centres in 26
Study Setting	countries world	· · ·
Participants		<i>eria</i> : Age = 18 years to 55 years, diagnosis of RRMS
i ai cicipantis		teria), EDSS = 0 to 5.0; with $\geq$ 1 relapse in the year
	before entry in	
	•	<i>criteria</i> : Those who had signs or symptoms of other
		IS; progressive forms of MS; heart disease; treat-
		ced or participated in the previous trials of drug for
		severe depression; alcohol or drug misuse; suicide
	•	us or acute live, renal, or bone marrow dysfunction;
		mmaglobulinopathy, or uncontrolled epilepsy; con-
		r allergy to the drug used in the study; unable to have
	MRI.	i unergy to the drug used in the study, unusie to have
		<u>acteristics in glatiramer acetate group:</u> median
		(27-43); 68% female; EDSS median (range) 2 (1,5-
	3,0) mean 2,28	
Intervention group		tate 20 mg SC q.d. (n = 448)
Comparison group		1-1b 250  mcg SC every other day  (n = 897)
		1-1b 500  mcg SC every other day  (n = 899)
Outcome	<u>Primary endpoints:</u> Relapse-based outcomes at year 2 (ARR, days	
	to first relapse, proportion relapse-free).	
	<u>Secondary endpoints:</u> Confirmed EDSS progression; MS-related	
		nospital, MS-related steroid course, different MRI
	outcomes.	
	Definitions use	d for endpoints: <u>Relapses:</u> New or recurrent neuro-
	logical abnormalities that were separated by at least 30 days from	
	the onset of the preceding event, lasted at least 24 hours, and oc-	
	curred without fever or infection.	
	<i>EDSS progression</i> : Measured as a 1-point change in the score that	
	was sustained for 3 months.	
Follow-up	2 to 3,5 years	
Treatment history	Treatment-naive (based on inclusion criteria).	
Critical appraisal		
Randomization		Adequate
Allocation concealmen	it	Adequate
Double-blinding		No [(rater-blinded), IFN doses double-blinded]
<b>Baseline characteristic</b>	e similarity	Yes
Outcome measures		Adequate
Withdrawals		15%
ITT Analysis		Unclear
Funding		Manufacturer

Calabrese et al., 2012 (47), included in Khai et al. (27)

		, ,	
<b>RCT identification</b>	Not reported		
Study setting	Rater-blinded, randomized controlled trial, single-centre in Italy		
Participants	<i><u>Eligibility criteria</u></i> : Age = 18 years to 55 years, diagnosis of RRMS		
	(McDonald/Polman criteria), EDSS = 0 to 5.0		
	Key exclusion criteria: Those previously treated with immunosup-		
	pressive drugs.		
	Baseline characteristics: Age: 37+/-10 years; 70% female; EDSS		
	2,0+/-1,1		
Intervention group	Glatiramer acetate 20 mg SC q.d. ( $n = 55$ )		
Comparison group	Interferon beta-1a 44 mcg SC t.i.w. $(n = 55)$		
	Interferon beta-1a 30 mcg IM q.w. (n = 55)		
Outcome	Different MRI outcomes.		
	Annualized relapse rate.		
	EDSS change.		
	Definition not stated		
Follow-up	2 years	2 years	
Treatment history	Unclear (inadequate information to characteristics)		
Comments		n also includes a group of disease modifying treated	
	patients, and disease modifying drug untreated controls		
Critical appraisal			
Randomization		Adequate	
Allocation concealment	ıt	Adequate	
Double-blinding		No (rater blinded)	
Baseline characteristic similarity		Yes	
Outcome measures		Adequate	
Withdrawals		15%	
ITT Analysis		No	
Funding		Manufacturer	

GALA (Glatiramer Acetate Low-frequency Administration) study, Khan et al., 2013 (35), not included in Khai et al. (27)

RCT identification	1	
	Not reported	11 11 1 . 1 . 1
Study setting	-	uble-blind study was conducted in 142 sites in 17
		ng the United States, Bulgaria, Croatia, Germany,
	Poland, Romania,	
Participants	<u>Eligibility criteric</u>	18 to 55 years of age, Confirmed RRMS diagno-
		he revised McDonald criteria), had an Expanded
		cale (EDSS) score of <=5.5, and were relapse-free
		Patients also were required to have >=1 docu-
		n the 12 months prior to screening, >=2 docu-
	mented relapses	in the 24 months prior to screening, or 1 docu-
	mented relapse be	etween 12 and 24 months prior to screening with
	at least 1 docume	nted T1 gadolinium enhancing lesion in an MRI
	performed within	12 months of screening.
	<i>Key exclusion crit</i>	teria: Several exclusions criteria based on previ-
	ous and/or concu	
	Baseline characte	eristics in placebo group: 38+/-9 years; 68% fe-
	male; EDSS 2.7+/	
Intervention group	Glatiramer acetat	e sc 40mg (1ml) tiw (n=943)
Comparison group	Placebo (n=461)	
Outcome	Primary endpoint: Annualised relapse rate	
		<u>ints:</u> MRI outcomes
	Definition used for	or relapse: A Relapse was defined as the appear-
	ance of $>=1$ new neurological abnormalities or the reappearance of	
	>=1 previously observed neurological abnormalities lasting at least	
	48 hours and preceded by an improving neurological state of at	
	least 30 days from the onset of previous relapse. An event was	
	counted as a relapse when the patient's symptoms were accompa-	
	nied by observed objective neurological changes consistent with an	
	increase of $\geq =0.5$ points in the EDSS score compared with previous	
	evaluation, or an increase of 1 grade in the actual score of $>=2$ or	
	more of the 7 FSs; or an increase of 2 grades in the score of 1 FS,	
		e previous assessment.
Follow-up	12 months (placeb	
Treatment history	Mixed (based on e	
Risk of bias		
Random sequence gen	eration	Low risk
Allocation concealmen		Not described, but blinding is adequate.
Blinding of participant and personnel		Low risk
Blinding of outcome as		Low risk
Incomplete outcome data		Low risk
-		Analysis performed as ITT
Selective reporting		Not detected
Other sources of bias		Funding: Manufacturer

CombiRx study 2013. Lublin et al., (48), included in Khai et al. (27)

RCT identification	NCT00211887			
Study setting				
Study setting	A double-blind, randomized, controlled study. 68 sites, both pri- vate practice and academic, in the USA and Canada			
Participants	<u>Eligibility criteria:</u> Patients with a diagnosis of RRMS by Poser or			
Farticipants	McDonald cirteria, aged 18- 60, EDSS score of 0 to 5.5, at least 2			
		0		
	exacerbations in the prior 3 years, where 1 exacerbation could be an magnetic resonance imaging (MRI) change meeting the 2001			
	McDonald MRI criteria for dissemination in time			
	<u>Key exclusion criteria:</u> prior history of seizure activity			
	Prior use of either interferon or glatiramer acetate			
	<i>Baseline characteristics:</i> Age: 38.0 +/- 10, 72% female, EDSS			
	<u>Baseline characteristics:</u> Age: 38.0 +/- 10, 72% female, EDSS score: 2.0 +/- 1.2			
Intervention group	Interferon beta-1a 30µg IM q.d and glatiramer acetate (GA) 20mg			
intervention group	q.d (n=499) (This group was outside our scope)			
	Glatiramer acetate 20mg q.d (n=259)			
	Interferon beta-1a 30µg IM q.w (n=250)			
Comparison group	Interventions were compared one with another			
Outcome	<b>≜</b>			
Outcome	<u>Primary endpoint:</u> Annualized relapse rate. <u>Secondary endpoints:</u> Disability progression (EDSS change or			
		, different MRI outcomes.		
	wore change)	, unterent with outcomes.		
	Definitions use	d for: <i>Relanses</i> : New or worsening neurologic symp-		
		<u>Definitions used for: <i>Relapses</i></u> : New or worsening neurologic symp- toms that lasted at least 24 hours without fever or infection, pre-		
	ceded by 30 days of stability.			
	<u>Disability progression:</u> 1.0 increase in the EDSS from baseline,			
	when baseline $\leq 5.0$ ; or an increase of 0.5 from baseline, when			
	baseline $\geq$ 5.5, sustained for 6 months (2 successive quarterly vis-			
	its), as assessed by the blinded EDSS examiner and confirmed cen-			
	trally.			
Follow-up	3 years			
Treatment history		ve (based on exclusion criteria)		
Critical appraisal	•			
Randomization		Adequate		
Allocation concealeme	ent	Adequate		
Double-blinding		Yes		
Baseline characteristi	c similarity	Yes		
Outcome measures	•	Adequate		
Withdrawals		18%		
ITT analysis		Yes		
Funding		Public, study agents and placebo provided by man-		
		ufacturer		

## Interferon beta 1a (im)

MSCRG (Multiple Sclerosis Collaborative Research Group) study 1996,
Jacobs et al. (49), included in Khai et al. (27)

RCT identification	Not reported		
Study setting		andomized controlled trial. 4 centres in the US	
Participants	<i><u>Eligibility criteria:</u> Age = 18 years to 55 years, diagnosis of relaps-</i>		
1 ai ticipants	ing MS (complete and incomplete remissions) (Poser et al.), EDSS		
	= 1 to 3.5; had $\geq$ 2 relapses in previous 3 years, no exacerbations for		
	at least 2 months at study entry		
	<u>Key exclusion criteria:</u> Prior immunosuppressant or IFN therapy;		
	adrenocorticotropic hormone or corticosteroid treatment with 2		
	months of entry; pregnancy or nursing; unwillingness to practice		
	contraception; presence of chronic-progressive MS, or any disease		
	other than MS compromising organ function.		
	Baseline characteristics: Age 37+/-7; 73% female; EDSS: 2.4+/-		
	0.8		
Intervention group	Interferon beta-l a 30 mcg IM q.w. (n=158)		
Comparison group	Placebo (n=143)		
Outcome	Primary endpoint: Time to onset of sustained worsening in disa-		
	bility.		
	Secondary endpoints: Proportion of patients with relapses, annu-		
	alized relapse r	ate, different MRI outcomes	
	Definitions	d for ordeninte. Delances. The encourse of new	
	<u>Definitions used for endpoints:</u> <u>Relapses:</u> The appearance of new		
	nourological st	motome or worsening of pre-existing neurological	
		mptoms or worsening of pre-existing neurological	
	symptoms lasti	ing at least 48 hours in a patient who had been neu-	
	symptoms lasti rologically stab	ing at least 48 hours in a patient who had been neu- le or improving for the previous 30 days, accompa-	
	symptoms lasti rologically stab nied by objecti	ing at least 48 hours in a patient who had been neu- le or improving for the previous 30 days, accompa- ve change on neurological examination.	
	symptoms lasti rologically stab nied by objecti <i>Disability prog</i>	ing at least 48 hours in a patient who had been neu- le or improving for the previous 30 days, accompa- ve change on neurological examination. <u>pression:</u> Deterioration from baseline by at least 1.0	
Follow-up	symptoms last rologically stab nied by objectiv <u>Disability prog</u> point on the EI	ing at least 48 hours in a patient who had been neu- le or improving for the previous 30 days, accompa- ve change on neurological examination.	
Follow-up Treatment history	symptoms lasti rologically stab nied by objectiv <u>Disability proc</u> point on the EI 2 years	ing at least 48 hours in a patient who had been neu- le or improving for the previous 30 days, accompa- ve change on neurological examination. <u>pression:</u> Deterioration from baseline by at least 1.0 DSS persisting for at least 6 months.	
Follow-up Treatment history	symptoms lasti rologically stab nied by objectiv <u>Disability proc</u> point on the EI 2 years	ing at least 48 hours in a patient who had been neu- le or improving for the previous 30 days, accompa- ve change on neurological examination. <u>pression:</u> Deterioration from baseline by at least 1.0 DSS persisting for at least 6 months.	
	symptoms lasti rologically stab nied by objectiv <u>Disability proc</u> point on the EI 2 years Treatment-naiv	ing at least 48 hours in a patient who had been neu- le or improving for the previous 30 days, accompa- ve change on neurological examination. <u>pression:</u> Deterioration from baseline by at least 1.0 DSS persisting for at least 6 months.	
Treatment history Critical appraisal Randomization	symptoms lasti rologically stab nied by objectiv <u>Disability proc</u> point on the EI 2 years Treatment-naiv clinical expert	ing at least 48 hours in a patient who had been neu- le or improving for the previous 30 days, accompa- ve change on neurological examination. <u>pression:</u> Deterioration from baseline by at least 1.0 DSS persisting for at least 6 months.	
Treatment history Critical appraisal Randomization Allocation concealment	symptoms lasti rologically stab nied by objectiv <u>Disability proc</u> point on the EI 2 years Treatment-naiv clinical expert	ing at least 48 hours in a patient who had been neu- le or improving for the previous 30 days, accompa- ve change on neurological examination. <u>pression:</u> Deterioration from baseline by at least 1.0 DSS persisting for at least 6 months. ve (based on exclusion criteria, year of study, and input).	
Treatment history Critical appraisal Randomization Allocation concealmen Double-blinding	symptoms lasti rologically stab nied by objecti <u>Disability prog</u> point on the EI 2 years Treatment-nai clinical expert i	Adequate Adequate Adequate Adequate Yes	
Treatment history Critical appraisal Randomization Allocation concealmen Double-blinding Baseline characteristic	symptoms lasti rologically stab nied by objecti <u>Disability prog</u> point on the EI 2 years Treatment-nai clinical expert i	ing at least 48 hours in a patient who had been neu- ble or improving for the previous 30 days, accompa- ve change on neurological examination. <u>mession:</u> Deterioration from baseline by at least 1.0 DSS persisting for at least 6 months. ve (based on exclusion criteria, year of study, and input). Adequate Adequate Yes Yes	
Treatment history Critical appraisal Randomization Allocation concealmen Double-blinding Baseline characteristic Outcome measures	symptoms lasti rologically stab nied by objecti <u>Disability prog</u> point on the EI 2 years Treatment-nai clinical expert i	ing at least 48 hours in a patient who had been neu- le or improving for the previous 30 days, accompa- ve change on neurological examination. <u>mession:</u> Deterioration from baseline by at least 1.0 DSS persisting for at least 6 months. ve (based on exclusion criteria, year of study, and input). Adequate Adequate Yes Yes Adequate	
Treatment history Critical appraisal Randomization Allocation concealmen Double-blinding Baseline characteristic Outcome measures Withdrawals	symptoms lasti rologically stab nied by objecti <u>Disability prog</u> point on the EI 2 years Treatment-nai clinical expert i	ing at least 48 hours in a patient who had been neu- le or improving for the previous 30 days, accompa- ve change on neurological examination. <u>mession:</u> Deterioration from baseline by at least 1.0 DSS persisting for at least 6 months. ve (based on exclusion criteria, year of study, and input). Adequate Adequate Yes Yes Yes Adequate 8%	
Treatment history Critical appraisal Randomization Allocation concealmen Double-blinding Baseline characteristic Outcome measures	symptoms lasti rologically stab nied by objecti <u>Disability prog</u> point on the EI 2 years Treatment-nai clinical expert i	ing at least 48 hours in a patient who had been neu- le or improving for the previous 30 days, accompa- ve change on neurological examination. <u>mession:</u> Deterioration from baseline by at least 1.0 DSS persisting for at least 6 months. ve (based on exclusion criteria, year of study, and input). Adequate Adequate Yes Yes Adequate	

# *EVIDENCE (EVidence of Interferon Dose-response: European North American Comparative Efficacy) study 2002, Panitch et al. (50), included in Khai et al. (27)*

ciuded în Khai et al. (2	1	1		
RCT identification	Not reported			
Study setting	Rater-blinded, randomized, placebo-controlled trial in 56 centres			
	in Europe, Canada, and US.			
Participants	<i><u>Eligibility criteria</u></i> : Age = 18 years to 55 years, IFN-naive patients			
	with definite RRMS (Poser et al.), EDSS = 0 to $5.5$ ; $\ge 2$ exacerba-			
	tions of MS in the prior 2 years.			
	Key exclusion criteria: use of defined treatments in previous peri-			
	ods.			
	Baseline characteristics in-30 mcg IM q.w group: Age 37,4 years			
		74,6%female, EDSS median 2,0 mean 2,3		
Intervention group	Interferon beta-1a 30 mcg IM q.w. (n = 338)			
	Interferon beta-	Interferon beta-1a 44 mcg SC t.i.w. (n = 339)		
Comparison group	These drugs were compared one with another			
Outcome	<u>Primary endpoint:</u> Proportion of patients who were relapse-free at			
	24 weeks.			
	Secondary endpoints: Relapse, disability, and MRI outcomes at 48			
	weeks.			
		<u>d for endpoints: <i>Relapses</i>:</u> The appearance of new		
	symptoms or worsening of an old symptom, accompanied by an ap-			
	propriate objective finding on neurologic examination by the			
	blinded evaluator, lasting at least 24 hours in the absence of fever			
	and preceded by at least 30 days of clinical stability or improve-			
		ment.		
	<u>Disability</u> : Progression by one point on the EDSS scale confirmed			
	at a visit 3 or 6 months later without an intervening EDSS value			
	that would not meet the criteria for progression.			
Follow-up	24 weeks (treatment for 24 weeks, follow-up until 48 weeks)			
Treatment history	Unclear (inadeq	uate information to characterise)		
Critical appraisal				
Randomization		Adequate		
Allocation concealme	nt	Adequate		
Double-blinding		No (rater-blinded)		
Baseline characteristic similarity		Yes		
Outcome measures		Adequate		
Withdrawals		4%		
ITT Analysis		Yes		
Funding		Manufacturer		
<b>v</b>				

INCOMIN (INdependent COMparison of INterferons) study, Durelli et al.2002,(51), included in Khai et al. (27)

<b>RCT identification</b>	Not reported	
Study setting	Open label, rate	er-masked, randomized controlled trial in 15 centres
	in Italy	
Participants		eria: Age = 18 years to 50 years, clinically definite
	RRMS (Poser e	t al.), EDSS = 1-3.5; had two clinically documented
	relapses during	the preceding 2 years, and no relapse (and no cor-
	ticosteroid trea	tment) for at least 30 days before the study entry.
	Key exclusion of	<u>eriteria:</u> Previous systemic treatment with IFN beta
		rith other immunosuppressive or immunomodula-
	<i>i</i> 0 i	ept corticosteroids);
		<u>cteristics:</u> Age 37+/-8; 65% female; EDSS 2,0+/-0,7
Intervention group		-1a 30 mcg IM q.w. (n = 92)
Comparison group		-1b 250 mcg SC every other day $(n = 96)$
Outcome		<u>pint:</u> Proportions of patients free from relapses dur-
	ing 24 months.	
		<i>points</i> : Annualized relapse rate, annualized treated
		coportion of patients free from sustained and con-
		sion from disability, EDSS score, time to sustained
	and confirmed	progression in disability.
	<u>Definitions used for endpoints:</u> <u>Relapses:</u> The occurrence of new	
		mptoms or worsening of an old one, with an objec-
		at least one point in Kurtzke Functional System
		at least 24 hours, without fever, and which followed
		ical stability or of improvement of at least 30 days.
	<u>Disability progression:</u> An increase in EDSS of at least 1 point sus-	
		ast 6 months and confirmed at the end of follow-up.
Follow-up	2 years	•
Treatment history	Treatment-naiv	ve (based on exclusion criteria).
Critical appraisal		
Randomization		Adequate
Allocation concealmer	ıt	Adequate
Double-blinding		No (rater-masked)
<b>Baseline characteristic</b>	e similarity	Yes
Outcome measures		Adequate
Withdrawals		16%
ITT Analysis		Yes
Funding		Public

Clanet et al., 2002 (52), included in Khai et al. (27)

RCT identification	Not reported	
Study setting	Randomized, d	louble-blind, dose-comparison study. 38 centers in
	Europe	
Participants	<u>Eligibility crite</u> of MS (Poser e definite MS; wi <u>Key exclusion o</u> tinuous deterio months, witho year); had a re nant or breastfo ideation, or se products within ucts for MS trea pressant therap	<u>tria:</u> Age = 18 years to 55 years, with a relapsing form t al.), EDSS = 2.0 to 5.5; had a clinical diagnosis of th ≥ 2 relapses within 3 years before randomization. <u>criteria:</u> Progressive forms of MS (defined as a con- bration in neurologic function during the previous 6 but superimposed relapses during the previous 1 lapse within 2 months before randomization; preg- eeding; with history of uncontrolled seizure, suicidal were depression; received treatment with IFN beta n 3 months of randomization; investigational prod- atment or non-MS indications; chronic immunosup- by or chronic steroid therapy. <u>ucteristics:</u> Age; 37+/-8; 68% female; EDSS: 3.6+/-
	1.0;	
Intervention group	,	-1a 30 mcg IM once weekly (n=402)
	Interferon beta	-1a 60 mcg IM once weekly N=(400)
Comparison group	The two doses	of Interferon beta-1a are compared one with another
Outcome	<u>Primary endpoint:</u> Disability progression.	
	Secondary endpoint: Relapse rate, annualized IV steroid use, per-	
	cent of patients with relapse-free, different MRI outcomes.	
	<u>Definitions used for endpoints:</u> <u>Relapses:</u> Not reported. <u>Disability progression:</u> Time to a sustained increase of ≥ 1.0 point on the EDSS persisting for 6 months for subjects with baseline EDSS scores ≤ 4.5, or a 0.5 point increase for subjects with a base- line EDSS score ≥ 5.0.	
Follow-up	At least 3 years	
Treatment history	Unclear (inade	quate information to characterise)
Critical appraisal		
Randomization		Insufficient reporting
Allocation concealmer	t	Insufficient reporting
Double-blinding		Yes
Baseline characteristic	e similarity	Yes
Outcome measures		Adequate
Withdrawals		30%
ITT Analysis		Yes
Funding		Manufacturer

Kappos et al., 2011 (36), included in Khai et al. (27)

RCT identification	NCT00676715		
Study setting		ontrolled study. 79 centres in 20 countries in North	
Study Setting		central Europe, Asia, western Europe, and Latin	
	America.	contrar Europe, rista, western Europe, and Eatin	
Participants		eria: Age = 18 years to 55 years, diagnosis of RRMS,	
i ul ticipulitis		had $\geq 2$ relapses in previous 3 years.	
		<i>criteria</i> : SPMS or PPMS, disease duration more than	
		ients with EDSS of 2 or less; history or presence of	
		ical systemic autoimmune disorders; treatment with	
		ymphocyte-depleting therapies; use of lymphocyte	
		trafficking disorders within previous 24 weeks; use of beta interfer- ons, glatiramer acetate, intravenous immunoglobulin, plasmapher-	
		nosuppressive treatments within previous 12 weeks,	
		glucocorticoids within previous 4 weeks; or intoler-	
	ance to IFN be	<b>.</b>	
		<u>icteristics in placebo group</u> : Age in years: 38 +/9,	
		ean EDSS score (-/+ SD): $3.2 +/- 1.4$	
Intervention group		00 mg IV day 1 and 15 ( $n=55$ , not our scope)	
inter tention group		00  mg IV day 1 and 15 (n = 55, not our scope)	
		a-1a 30 mcg IM q.d. (n=55)	
Comparison group	Placebo (n=54		
Outcome	1 0 1	<u>pint:</u> MRI outcomes.	
		<i>lpoints</i> : Annualized relapse rate, proportion of re-	
	lapse-free patients.		
	1 1		
	Definitions use	ed for endpoints: <u>Relapses:</u> The occurrence of new or	
	worsening neu	rological symptoms attributable to MS, and imme-	
	diately precede	ed by a stable or improving neurological state of at	
	least 30 days.		
	<u>Disability progression:</u> An increase of 1 point or more from base-		
		re confirmed at the next scheduled examination 3	
		nitial screening.	
Follow-up	24 weeks		
	· 1 /	xs, but after 24 weeks, comparator groups switched	
	to ocrelizumab		
Treatment history	Mixed (based of	on reported baseline characteristics)	
Critical appraisal			
Randomization	_	Insufficient reporting	
Allocation concealmen	nt	Not reporting	
Double-blinding		No	
	• •1 •.		
Baseline characteristi	c similarity	No	
Baseline characteristic Outcome measures	e similarity	No Adequate	
Baseline characteristic Outcome measures Withdrawals	e similarity	No Adequate 6%	
Baseline characteristic Outcome measures	e similarity	No Adequate	

Mokhber et al., 2014 (53), not included in Khai et al. (27)

<b>RCT identification</b>	Protocol numb	er: 84393-1
Study setting		andomized trial, single center in Iran
Participants	<u>Eligibility criteria</u> : Eligible participants were all new cases of definite MS according to the revised McDonald criteria, which include magnetic resonance imaging, detailed neurological history and examination, and paraclinical laboratory tests of cerebrospinal fluid findings and visual-evoked potential <u>Key exclusion criteria</u> : Patients were excluded if they had a history of substance abuse or prior treatment with any type of DMTs <u>Baseline characteristics</u> : Age 29,+/-8; 65% female; EDSS: mean=2.02	
Intervention group		a-1a (Avonex ) 30 mcg once per week IM injection;
	(n=23) Interferon beta-1a (Rebif) 44 mcg t.i.w. SC injection; (n=23) Interferon beta-1a (Betaferon) 0.25 mg every other day SC injection (n=23)	
Comparison group	These drugs were compared one with another	
Outcome		<u>pint:</u> Cognition status I <u>point:</u> EDSS scale
Follow-up	1 year	
Treatment history	Treatment-naive	
	Risk of Bias	
Random sequence gen	Adequate "The study neurologist (MRA) enr the participants and allocated the subjects us computer-generated list of random numbers"	
Allocation concealmen		
Blinding of participant nel	•	Assessors: yes Participants: insufficient reporting
Blinding of outcome as		Adequate
Incomplete outcome d	ata	6% lost to follow-up Modified analysis based on available data
Selective reporting		None detected
Other sources of bias		No conflict of interest declared. Funding seem to be public "The study was supported by the Vice Chancellor of Research at Mashhad University of Medical Sciences in Iran (Grant number:84393)"

### BRAVO (Benefit-Risk Assessment of AVonex and LaquinimOd ) study, Vollmer 2014 (54), not included in Khai et al. (27)

<b>RCT identification</b>	NCT00605215	
Study setting	-	bebo-controlled phase III trial in 155 sites in 18
	countries (including	g. USA and several European countries)
Participants	Eligibility criteria:	age 18–55 years, diagnosis of RRMS (revised
-		, and EDSS scores of $0-5.5$ . At least one relapse
	-	nonths, two in the previous 24 months, or one in
		months, plus one gadolinium-enhancing (GdE)
	lesion in the previo	
		<i>ria:</i> progressive forms of MS; use of glatiramer
		vious 2 months; and prior use of natalizumab,
		bine, or any interferon beta at any time.
	<b>1</b>	
		istics (in placebo group): Age (median and 25-75
	-	9,3-45,4); 71,3% female; EDSS (median and 25-75
-	percentile) 2.5 (1.5,	
Intervention group		g capsule q.d. (n=434)[not our scope]
		IM 30 mcg once-weekly injection ( $n = 447$ )
Comparison group		laquinimod) (n = 450)
Outcome	Primary endpoints	<u>:</u> Annualized relapse rate (ARR)
	Secondary endpoir	<i>its:</i> percent change in normalized brain volume
	from baseline to 2	4 months; changes in disability measured with
		ISFC z-score at 24 months/early termination)
		<i>ints:</i> confirmed worsening of EDSS scores sus-
		s. MRI endpoints: the cumulative numbers at 12,
		esions and of new or enlarging ([50 % larger than
	previous scan) T2 le	
		<u>r endpoints: <i>Relapse</i></u> = appearance of one or more
		phormalities, or reappearance of one or more pre-
		eurological abnormalities, in the absence of fever,
		B h, preceded by > 30 days of a stable or improving
		ompanied by at least one of the following: an in-
	crease of at least 0.5 point in EDSS score, an increase of one grade in	
		he seven functional systems (FS) on the EDSS, or
	an increase of two g	
		ion: a 1.0 point EDSS increase in EDSS if baseline
		5 if baseline score was 5.5, for 3 months.
Follow-up	2 years	
Treatment history	Mixed (based on ex	clusion criteria)
Risk of bias		
Random sequence ge		Low risk
Allocation concealme	nt	Not described. (Assume low risk based on description of sequence generation and blinding)
Blinding of participa	nt and personnel	Not for our comparison
Blinding of outcome		Adequate
Incomplete outcome		Low risk
	uală	None detected
Selective reporting		
Other sources of bias	Differences in mean T2 lesion volume and GdE lesions at baseline between laquinimod or IFNb-1a groups	

# Interferon beta 1a (sc)

# PRISMS (Prevention of Relapses and Disability by Interferon _beta 1a Subcutaneously in Multiple Sclerosis) study1998 (55), in Khai et al. (27)

-		
Not reported		
Double-blind, randomized, controlled trial. 22 centres in 9 co	un-	
tries including Australia, Canada, and Europe.	<u>.</u>	
	= 0	
to 5.0; had $\geq$ 2 relapses in previous 2 years.		
Key exclusion criteria: Previous systemic treatment with IFN, by	/m-	
modulatory or immunosuppressive treatments in the preceding	g 12	
	29-	
0		
	ac-	
tivity under MRI and burden of disease.		
	<u>Disability progression:</u> An increase in EDSS of at least 1 point sus-	
tained over at least 3 months.		
2 years		
Treatment-naive (based on exclusion criteria, year of study,		
Treatment-naive (based on exclusion criteria, year of study, clinical expert input).		
Treatment-naive (based on exclusion criteria, year of study, a clinical expert input). Adequate		
Treatment-naive (based on exclusion criteria, year of study, a clinical expert input).         Adequate         Adequate		
Treatment-naive (based on exclusion criteria, year of study, sclinical expert input). Adequate Adequate Yes		
Treatment-naive (based on exclusion criteria, year of study, a clinical expert input).         Adequate         Adequate         Yes         c similarity       Yes		
Treatment-naive (based on exclusion criteria, year of study, a clinical expert input).         Adequate         Adequate         Yes         c similarity       Yes         Adequate		
Treatment-naive (based on exclusion criteria, year of study, sclinical expert input).          Adequate         Adequate         Yes         c similarity       Yes         Adequate         10%		
Treatment-naive (based on exclusion criteria, year of study, a clinical expert input).         Adequate         Adequate         Yes         c similarity       Yes         Adequate		
	Double-blind, randomized, controlled trial. 22 centres in 9 co tries including Australia, Canada, and Europe. <u>Eligibility criteria</u> : Adult RRMS patients (Poser et al.), EDSS to 5.0; had $\geq$ 2 relapses in previous 2 years. <u>Key exclusion criteria</u> : Previous systemic treatment with IFN, by phoid irradiation, or cyclophosphamide, or with other immu modulatory or immunosuppressive treatments in the preceding months. <u>Baseline characteristics</u> : Age: median (interquartile range) 35 ( 40); 69% female; EDSS:2.5+/-1.2 Interferon beta-1a 22 mcg SC t.i.w. (n=189) Interferon beta-1a 44 mcg SC t.i.w. (n=184) Placebo (n=187) <u>Primary endpoints</u> : Times to first and second relapse, prop tion of relapse-free patients, disability progression, ambulation dex, need for steroid therapy and hospitalization, and disease tivity under MRI and burden of disease. <u>Definitions used for endpoints</u> : <u>Relapses</u> : The appearance of a r symptom or worsening of an old symptom over at least 24 ho that could be attributed to MS activity and was preceded by stabi or improvement for at least 30 days.	

*IMPROVE (Investigating MRI Parameters with RebifimprOVEd formulation) study 2010, De Stefano et al., (94), included in Khai et al. (27)* 

<b>RCT identification</b>	NCT00441103	
Study setting	Double-blind,	randomized, placebo-controlled trial, multi-centre,
	multi-country i	in European countries.
Participants	Eligibility crite	eria: Age = 18 years to 60 years, diagnosis of RRMS
		teria), EDSS = 0 to 5.5; active disease ( $\geq$ 1 clinical
		gadolinium-enhancing MRI lesion) within the 6
		before randomization.
	<u>Exclusion criteria</u> : Not specified.	
		<u>cteristics:</u> Not reported
Intervention group		-1a 44 mcg SC t.i.w. (n = 120)
Comparison group	Placebo ( $n = 60$	
Outcome		<u>pint:</u> Number of combined unique active MRI brain
	lesions at week	
		dpoints: Number of combined unique active le-
		scan, other MRI outcomes, relapse rate.
Follow-up	16 weeks	
Treatment history	Unclear (inadequate information to characterise)	
Comments	Double-blind phase:16 weeks. After that, patients received Inter-	
	feron beta-1a, 44 mg sc tiw, for 24 weeks (rater-blind phase).	
	The analysis populations for the rater-blind period comprised pa-	
		pleted treatment during the double-blind period
	(Interferon bet	a-1a, n=12; placebo,n=57).
Critical appraisal		
Randomization	•	Insufficient reporting
Allocation concealmen	t	Not reporting
Double-blinding		Yes
Baseline characteristic	similarity	Not reporting
Outcome measures		Adequate
Withdrawals		Not reporting
ITT Analysis		Yes
Funding		Manufacturer

# Interferon beta 1b (sc) IFNB-MS 1993, (57), included in Khai et al. (27)

<b>RCT identification</b>	Not reported	
Study setting		lacebo-controlled trial Multi-centre Canada and the
	US.	
Participants	Eligibility crite	<i>ria</i> : Age = 18 years to 50 years, diagnosis of RRMS
-	(McDonald crit	teria), EDSS = 0 to 5.5; had $\geq$ 2 exacerbations during
		years; clinically stable for at least 30 days before en-
		d no adrenocorticotrophic hormone or prednisone
	during this per	
		<i>criteria</i> : Prior treatment with azathioprine or cyclo-
	phosphamide.	
		<u>cteristics:</u> Age 35+/-7; 70% female; EDSS 2,9+/-1,1
Intervention group		-1b 250  mcg SC every other day (n = 124)
~ .		-1b 50 mcg SC every other day (n=125)
Comparison group	Placebo (n = 12	
Outcome		<u>pints:</u> Annualized relapse rate, proportion of relapse-
	free patients	having Time to first volge as here denotion and
		<i>points:</i> Time to first relapse, relapse duration and
	severity, change	e in EDSS, MRI outcomes.
	Definitions use	ed for endpoints: <i>Relapses</i> : The appearance of a new
	symptoms or worsening of an old symptom, attributable to MS; ac-	
	companied by an appropriate new neurologic abnormality; lasting	
		rs in the absence of fever; and preceded by stability
		it for at least 30 days.
		<i>ression</i> : A patient was considered to have progres-
	sion in disability when there was a persistent increase of 1 or more	
	EDSS points confirmed on two consecutive evaluations separated	
	by at least 3 months.	
Follow-up	3 years	
Treatment history	Treatment-naiv	ve (based on year of study and clinical expert input).
Critical appraisal		
Randomization		Insufficient reporting
Allocation concealmen	nt	Not reporting
Double-blinding		Yes
Baseline characteristic	c similarity	Yes
Outcome measures		Adequate
Withdrawals		33%
ITT Analysis		Yes
Funding		Not reporting

# Etemadifar et al., 2006(58), included in Khai et al. (27)

<b>RCT identification</b>	Not reported	
Study setting	Rater-blinded, r	andomized controlled trial, neurology outpatient
	clinics in Iran	
Participants		<i>ia</i> : Age = 15 years to 50 years, diagnosis of relaps-
		t al.), EDSS = 0 to 5.0; $\geq$ 2 relapses within the 2-
		eatment initiation documented by a neurologist.
		<i>iteria</i> : History of severe allergic or anaphylactic re-
	action to any IFN, or to other components of drug formulation; ev-	
	idence of neurologic, psychiatric, cardiac, endocrinologic, hemato-	
		enal, active malignancy, autoimmune diseases, or
		sease; history of uncontrolled seizure or suicidal
		e depression; lactation and pregnancy.
		<u>eristics:</u> Age 29+/-7; 76% female; EDSS 2,0+/-0,9
Intervention group		1b 250 mcg SC every other day $(n = 30)$
		1a 30 mcg IM q.w. (n = 30)
		1a 44 mcg SC t.i.w. (n = 30)
Comparison group		e compared one with another
Outcome	<u>Endpoints</u> : Number of relapses, proportion of relapse-free pa-	
	tients, EDSS scores	
	Definitions used for endpoints: <u>Relapses:</u> The appearance of a new	
	neurologic symptom, or severe deterioration in a pre-existing	
	symptom that lasted 24 hours causing the deterioration in the EDSS with 1 point.	
Follow-up	2 years	
Treatment history		uate information to characterise)
Critical appraisal	Unclear (madey	uate information to characterise)
Randomization	1	Insufficient reporting
Allocation concealmen		Not reporting
Double-blinding		No (rater-blinded)
Baseline characteristic		No
Outcome measures		Adequate
Withdrawals		0%
ITT Analysis		
Funding	Yes Not reporting	

### Natalizumab

### AFFIRM (Natalizumab Safety and Efficacy in Relapsing Remitting Multiple Sclerosis) study, Polman et al., 2006 (60), in Khai et al. (27)

<b>RCT identification</b>	NCT00027300	)	
Study setting	Randomized, d	louble-blind, placebo-controlled trial in 99 centres	
	in Europe, Nor	th America, Australia, and New Zealand.	
Participants		eria: Age = 18 years to 50 years, diagnosis of RRMS	
		teria), EDSS = 0 to 5.0; had MRI lesions with MS,	
		ly documented relapse within 12 months before the	
	study began.		
		<i>criteria</i> : relapse within 50 days before administra-	
		st dose of the study drug; treatment with specific	
		aceuticals (MS related)	
	Baseline characteristics: Age 36+/-8 years; 70% female; EDSS		
	2,3+/-1,2		
Intervention group		00 mg IV every 4 weeks (n = 627)	
Comparison group	Placebo (n = $3$		
Outcome		oints: Rate of clinical relapse at 1 year; cumulative	
		sustained progression of disability at 2 years.	
		<i>lpoints</i> : Different MRI outcomes at 1 and 2 years;	
		elapse-free patients at 1 year; progression of disabil-	
		neasured by MSFC.	
		<u>bints:</u> HRQoL was assessed by SF-36 (PCS and MCS)	
	and Subject Gl	obal Assessment Visual Analogue Scale.	
	Definitions	od for ondpoints: Palancac: Now or requirement nouro	
	<u>Definitions used for endpoints:</u> <u>Relapses:</u> New or recurrent neuro- logic symptoms not associated with fever or infection that lasted for		
	at least 24 hours and were accompanied by new neurologic signs		
	found by the examining neurologist.		
	Sustained progression of disability: An increase of 1.0 or more on		
	the EDSS from a baseline score of 1.0 or more or an increase of 1.5		
		a baseline score of 0 that was sustained for 12 weeks	
		build not be confirmed during a relapse).	
Follow-up	2 years		
Treatment history		quate information to characterise)	
Critical appraisal	eneren (muue		
Randomization		Adequate	
Allocation concealment	nt	Adequate	
Double-blinding		Yes	
<b>Baseline characteristic</b>	c similarity	Yes	
Outcome measures	•	Adequate	
Withdrawals		9%	
ITT Analysis		Yes	
Funding		Manufacturer	

Gobbi et al. (31), not included in Khai et al. (27)

RCT identification	NCT011	44052
Study setting		nized controlled study, rater blinded. One centre, Switzer-
Study Setting	land.	mzed controlled study, later billided. One centre, Switzer-
Donticinente		ity anitania, Datianta with DDMS (2005 MaDanald's anita
Participants		ity criteria: Patients with RRMS (2005 McDonald's crite-
		ed between 18 and 60 years, who were on natalizumab
		and feared or were at significant risk for progressive multi-
		ucoencephalopathy (PML) [Risk for PML was defined sig-
		in case of NTZ treatment duration equal to or greater than
		nths]. Patients had to be free of disease activity while
	on NTZ (free from relapses and disability progression for at least 6	
		and no gadolinium enhancing lesions on baseline MRI
		lusion criteria: relevant neurologic, internistic or psychiat-
		orders; treatment with steroids less than 1 month before
		ntry; treatment with any immunomodulators or immune-
		ssors other than steroids, ACTH* or NTZ in the past year.
		<u>e characterics in NTZ group:</u> Age median (range): 43 (20-
		% female, EDSS score (median (range)): 3 (1.5-3.5)
Intervention group		ie on natalizumab 300 mg IV q.m. (n=10)
Comparison group	Switch to interferon beta-1b 250 mcg every other day (n=9)	
Outcome		<u>y endpoint</u> was time to first on-study relapse
		ndomization.
	<u>Second</u>	<i>ary endpoints</i> included number of relapses, proportion of
	relapse free patients, severity of relapses (severe relapse was de-	
	fined by	$V \ge 1.5$ increase in EDSS score), 3 months confirmed disa-
	bility pi	cogression (defined by ≥1.0 increase in EDSS score), num-
	ber of n	ew T2-hyperintense lesions (nT2L) and Gd+L per patient
	at months 3, 6, 9 and 12.	
Follow-up	1 year	
Treatment history	Treatm	ent experienced
Risk of Bias		
Random sequence gen	era-	Adequate
tion		A monitoring agency prepared the randomization list and
		provided sealed envelopes for treatment allocation.
Allocation concealmen		
Blinding of participant		No
personnel		Rater blinded
Blinding of outcome as	ssess-	Adequate
ment		"EDSS and relapses assessment was performed by an ex-
		amining neurologist blinded to treatment."
Incomplete outcome d	ata	Analysis was based on intention to treat.
		Withdrawals: 10.5%
Selective reporting		None detected
Other sources of bias		Several of the authors report funding from one or several
		pharmaceutical companies.
* A OTTIL this shhused ation a		prairie of the second

*ACTH: this abbreviation was not explained in the publication

RESTORE-study 2014, Fox et al., (61), not included in Khai et al. (27)

RESIORE-Study 2014,		
<b>RCT identification</b>	NCT01071083	
Study setting	Randomized. parti	ally placebo-controlled study. 31 sites in North
2 8	America and Europ	
Participants		: Patients with RRMS receiving natalizumab,
i ai ticipants		
		ars, who had been treated with natalizumab for
		prior to randomization and who had no relapses
	during those 12 mc	
	Key exclusion crite	<i>ria:</i> presence of gadolinium enhancing lesions;
	presence of antinat	alizumab antibodies; immunosuppressive treat-
	ment within 24 mc	onths prior to randomization; treatment with IV
		plasmapheresis, or cytapheresis within 12
	0	ndomization; or treatment with systemic corti-
	-	months prior to randomization.
		<i>istics in placebo group</i> : Age: 40 +/- 10; 74% fe-
	male; EDSS: 3.3 +/	
Intervention group		ng IV every 4 weeks (n=45)
		modulatory therapy (IM interferon b-1a, glati-
		nethylprednisolone (n=88) [not included as pa-
	-	irologist selected the immunomodulatory ther-
		al basis; as such, the distribution of patients re-
	-	a, GA, and MP was not randomized, and the
	groups were unbal	
Comparison group	Placebo IV every 4	weeks (n=42)
Outcome	Relapse	
	Quality of life'	
		_
	Withdrawal dije to	adverse events
	Withdrawal due to	adverse events
	Withdrawal due to Deaths	adverse events
	Deaths	
	Deaths Definition used: Ra	adiographic and clinical disease activity. Quality
	Deaths Definition used: Ra of life with Visual	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact
	Deaths Definition used: Ra of life with Visual Scale, and cognitio	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis-
	Deaths Definition used: Ra of life with Visual	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis-
	Deaths Definition used: Ra of life with Visual Scale, and cognitio	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis-
Follow-up	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis-
Follow-up	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis- with EDSS.
	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression 24 weeks (52 week infusions of nataliz	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis- with EDSS. as but at week 28, patients resumed open-label umab)
Treatment history	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression 24 weeks (52 week infusions of nataliz	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis- with EDSS.
Treatment history Risk of Bias	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression 24 weeks (52 week infusions of nataliz Treatment experies	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis- with EDSS. as but at week 28, patients resumed open-label umab) need ( all groups received natalizumab at day 0)
Treatment history Risk of Bias Random sequence gen	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression 24 weeks (52 week infusions of nataliz Treatment experies	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis- with EDSS. as but at week 28, patients resumed open-label umab) nced ( all groups received natalizumab at day 0) Adequate
Treatment history Risk of Bias	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression 24 weeks (52 week infusions of nataliz Treatment experies	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis- with EDSS. as but at week 28, patients resumed open-label umab) aced ( all groups received natalizumab at day 0) Adequate Adequate
Treatment history Risk of Bias Random sequence gen Allocation concealmen	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression 24 weeks (52 week infusions of nataliz Treatment experies eration	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis- with EDSS. as but at week 28, patients resumed open-label umab) aced ( all groups received natalizumab at day 0) Adequate Adequate For arms natalizumab + placebo
Treatment history Risk of Bias Random sequence gen	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression 24 weeks (52 week infusions of nataliz Treatment experies eration	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis- with EDSS. as but at week 28, patients resumed open-label umab) need ( all groups received natalizumab at day 0) Adequate Adequate For arms natalizumab + placebo Adequate
Treatment history Risk of Bias Random sequence gen Allocation concealmen	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression 24 weeks (52 week infusions of nataliz Treatment experies eration	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis- with EDSS. as but at week 28, patients resumed open-label umab) aced ( all groups received natalizumab at day 0) Adequate Adequate For arms natalizumab + placebo
Treatment history Risk of Bias Random sequence gen Allocation concealmen Blinding of participan	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression 24 weeks (52 week infusions of nataliz Treatment experies eration at t and personnel	Adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis- with EDSS. as but at week 28, patients resumed open-label umab) need ( all groups received natalizumab at day 0) Adequate For arms natalizumab + placebo Adequate For arms natalizumab + placebo
Treatment history Risk of Bias Random sequence gen Allocation concealmen	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression 24 weeks (52 week infusions of nataliz Treatment experies eration at t and personnel	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis- with EDSS. as but at week 28, patients resumed open-label umab) nced ( all groups received natalizumab at day 0) Adequate For arms natalizumab + placebo Adequate For arms natalizumab + placebo Adequate For arms natalizumab + placebo Adequate
Treatment history Risk of Bias Random sequence gen Allocation concealmen Blinding of participan Blinding of outcome a	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression 24 weeks (52 week infusions of nataliz Treatment experies ceration nt t and personnel ssessment	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis- with EDSS. as but at week 28, patients resumed open-label umab) need ( all groups received natalizumab at day 0) Adequate For arms natalizumab + placebo Adequate For arms natalizumab + placebo Adequate For arms natalizumab + placebo Adequate For arms natalizumab + placebo
Treatment history Risk of Bias Random sequence gen Allocation concealmen Blinding of participan Blinding of outcome a Incomplete outcome d	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression 24 weeks (52 week infusions of nataliz Treatment experies ceration nt t and personnel ssessment	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis- with EDSS. as but at week 28, patients resumed open-label umab) need ( all groups received natalizumab at day 0) Adequate For arms natalizumab + placebo Adequate For arms natalizumab + placebo Adequate
Treatment history Risk of Bias Random sequence gen Allocation concealmen Blinding of participan Blinding of outcome a	Deaths Definition used: Ra of life with Visual Scale, and cognitio ability progression 24 weeks (52 week infusions of nataliz Treatment experies ceration nt t and personnel ssessment	adiographic and clinical disease activity. Quality Analogue Scale, and Modified Fatigue Impact n (Symbol Digit Modalities Test (SDMT)). Dis- with EDSS. as but at week 28, patients resumed open-label umab) need ( all groups received natalizumab at day 0) Adequate For arms natalizumab + placebo Adequate For arms natalizumab + placebo Adequate For arms natalizumab + placebo Adequate For arms natalizumab + placebo

Zecca et al., 2014 (32), not included in Khai et al. (27)

<b>RCT identification</b>	NCT1144052,							
Study setting	Randomized, rater-	blinded, parallel-group study, single center,						
	Switzerland							
Participants	Eligibility criteria: A	<i><u>Eligibility criteria</u></i> : Age between 18 and 60, being at significant risk						
		nt duration equal to or greater than 12 months)						
		being free of disease activity (free from relapses						
		ession for at least 6 months and no gadolinium						
		d + L] on baseline [BL] MRI). RRMS accord-						
		ld criteria [13] from 2010 to 2011						
		tics in Interferon group: Mean (range) 39 (24-						
_		(9); EDSS median (range) 3,0 (1,5-3,5)						
Intervention group		ab monthly intravenous (i.v.) 300 mg (n=10)						
Comparison group		eron beta 1b subcutaneous (s.c.) 250 mcg every						
	other day (n=9)							
Outcome		ent of patients included Paced Auditory Serial						
		(PASAT), Fatigue Scale for Motor and Cogni-						
		C), Functional Assessment of Multiple Sclerosis						
E llesse ser		uol visual analogue scale (EQ-VAS)						
Follow-up	1 year							
Treatment history	-	ed (All patients previously treated with natali-						
Risk of bias	zumab)							
	anation	Unclean/Not described						
Random sequence gen Allocation concealmer		Unclear/Not described Unclear/Not described						
Blinding of participan		No						
Blinding of outcome as		Adequate (rater-blinded)						
Incomplete outcome d	ลเล	No 17/19 completed study (reasons listed) None detected						
Selective reporting Other sources of bias								
other sources of blas		Some of the authors have received compen- sation from one or several of pharmaceutical						
		-						
	companies							

# Peg-interferon

# ADVANCE study 2014, Calabresi et al., (59), not in Khai et al. (27)

RCT identification	NCT00906							
Study setting		nd, randomized controlled study. 183 neurology prac-						
		countries, including north and south America, Europe,						
	India							
Participants		criteria: diagnosis of relapsing-remitting multiple scle-						
		osis as defined by the McDonald criteria, aged 18–65 years, a						
		S score of 0–5 , and at least two clinically documented re-						
		ne previous 3 years, with at least one having occurred						
		past 12 months.						
	Key exclus	<i>ion criteria</i> : pre-specified laboratory abnormalities, and						
	previous ti	reatment with interferon for multiple sclerosis for more						
	than 4 wee	ks or discontinuation less than 6 months before baseline						
	Baseline ci	haracteristics in placebo group: Age: 36+/- 10; 72% fe-						
	male; EDS	S: 2.4 +/-1.2						
Intervention group	vention group Peg-interferon beta-1a 125 mcg SC once every 2 weeks (n=512)							
	Peg-interferon beta-1a 125 mcg SC once every 4 weeks (n=500)							
Comparison group	Placebo (n=500)							
Outcome	<u>Primary endpoints:</u> Annualised relapse rate at week 48, based on							
	number of relapses.							
		endpoints: The number of new or newly enlarging hy-						
		lesions on T2-weighted images(relative to baseline						
		portion of patients who relapsed, and proportion of pa-						
		disability progression at 48 weeks.						
		<i>ndpoints:</i> Prespecified MRI endpoints at 48 weeks						
Follow-up		t placebo controlled only for 48 weeks						
Treatment history								
Risk of bias	Mixed (bas	sed on exclusion criteria)						
	<b></b>	V						
Random sequence gen		Yes						
Allocation concealment	it	Adequate						
		Patients received either study drug or placebo every 2						
		weeks to maintain masking; those assigned to receive						
		study drug every 4 weeks received alternate injections						
		of placebo and peg-interferon beta-1a every 2 weeks						
Blinding of participant	t and per-	Adequate "						
sonnel								
Blinding of outcome assess- Adequate								
ment								
Incomplete outcome d	ata	Adequate						
-	Intention to treat							
Selective reporting		None detected						
Other sources of bias		Funding: manufacturer						

# Teriflunomide

# O'connor et al., 2006 (62), included in Khai et al. (27)

RCT identification	Not reported							
Study setting		Randomized controlled study, double-blind. Centres in Canada						
Participants		<u>eria:</u> Age = 18 years to 65 years, with RRMS ( $n = 157$ )						
-		progressive MS with relapses $(n = 22)$ (Poser et al.),						
	EDSS = 0 to 6.	$DSS = 0$ to 6.0; had $\geq 2$ documented relapses in previous 3 yea						
	and one clinica	ne clinical relapse during the preceding year.						
		criteria: Prior treatment with interferon, gamma-						
		amer, or other non-corticosteroid immune-modula-						
		in the 4 months prior to the trial.						
		cteristics: Age: 39 +/-; 74% female; , EDSS score:						
	(median) 2.3							
Intervention group		oral 7 mg q.d. $(n=61)$						
		oral 14 mg q.d.(n=57)						
Comparison group	Placebo (n=61)							
Outcome	Primary endp	oint: Number of combined unique active (new and						
		ons per MRI scan during 36 weeks.						
		<i>lpoints:</i> Other MRI outcomes, number of patients						
		lapses, annualized relapse rate, number of relapsing						
	patients requir	red a course of steroids, EDSS change.						
	Definition use	d for Palance: The ennearance of a new symptom						
		<u>d for: <i>Relapses</i></u> : The appearance of a new symptom f an old symptom due to MS lasting 48 hours in the						
		er, preceded by period of stability of at least 30 days						
		ied by appropriate changes on neurologic examina-						
	tion.	ied by appropriate changes on neurologie examina						
Follow-up	36 weeks							
Treatment history	•	ve (based in exclusion criteria, year of study, and						
	clinical expert							
Comments		9% RRMS, 13.1% secondary progressive						
Critical appraisal								
Randomization		Insufficient reporting						
		Insufficient reporting						
Allocation concealme	nt	Insufficient reporting Not reporting						
Allocation concealmer Double-blinding	nt	1 U						
		Not reporting						
Double-blinding		Not reporting Yes						
Double-blinding Baseline characteristi		Not reporting Yes Yes						
Double-blinding Baseline characteristi Outcome measures		Not reporting       Yes       Yes       Adequate						

RCT identification	NCT00134563						
Study setting		randomized controlled trial. 127 centres in 21 coun-					
3		Canada, Europe, and US.					
Participants		<i>ria</i> : Age = 18 years to 55 years; diagnosis of RRMS					
-	(McDonald crit	teria), EDSS = 0 to 5.5; had $\geq$ 2 relapses in the pre-					
		ous 2 years or $\geq$ 1 relapse during the preceding year, but no re					
		days before randomization.					
		criteria: Had other systemic diseases; pregnant, or					
	planned to con	ceive during the trial period.					
		cteristics: Age 38+/-9; 72% female; EDSS: 2.7+/-1.3					
Intervention group	Teriflunomide	Teriflunomide oral 7 mg q.d. (n=365)					
	Teriflunomide	oral 14 mg q.d. (n=358)					
Comparison group	Placebo (n=36						
Outcome		<u>pint:</u> Annualized relapse rate.					
		lpoints: Disability progression (EDSS change), dif-					
	ferent MRI out	comes.					
		ed for endpoints: <u>Relapses:</u> The appearance of a new					
		symptom, or clinical worsening of a previous sign or					
		had been stable for at least 30 days and that persisted					
		of 24 hours in the absence of fever.					
		gression: An increase from baseline of at least 1.0					
		OSS score (or at least 0.5 points for patients with a					
		score greater than 5.5) that persisted for at least 12					
	weeks.						
Follow-up	108 weeks						
Treatment history	Mixed (based o	on reported baseline characteristics)					
Critical appraisal							
Randomization	_	Adequate					
Allocation concealment	nt	Adequate					
Double-blinding	• •1 •-	Yes					
Baseline characteristi	c similarity	Yes					
Outcome measures     Adequate							
Withdrawals     27%							
ITT Analysis		Yes					
Funding     Manufacturer							

TOWER-(Teriflunomide Oral in people With relapsing multiplE sclerosis) study, Confavreux et al. 2014 (65), not included in Khai et al. (27)

<b>RCT identification</b>	NCT00751881						
Study setting		uble-blind, placebo-controlled in 189 centres					
Study setting							
Donticinanta		ased sites in 26 countries					
Participants		a: ambulatory patients with RMS, aged 18–55					
		scores $\langle =5.5 \text{ and } \rangle =1$ relapse in the previous 12					
		lapses in the prior 24 months					
		teria: previously or concomitantly received cyto-					
		rferon beta, or glatiramer acetate within 3 months					
	-	, or had ever used natalizumab or other immuno-					
	suppressive agent						
		eristics (in placebo group): Age: 38+/-9; 70% fe-					
	male; EDSS: 2,7+						
Intervention group		mg once daily $(n=372)$					
	Teriflunomide 7 n	ng once daily (n=408)					
Comparison group	Placebo once dail	y (n=389)					
Outcome	Primary endpoin	ts: Annualised relapse rate (number of relapses					
	per patient-year)						
	Secondary endpo	<i>vints:</i> time to 12 week sustained accumulation of					
	disability; time to	fi rst relapse, proportion of patients free from re-					
	lapses, proportion	of patients free of accumulation of disability, and					
	change from base	line in EDSS score at week 48, and change in Fa-					
	tigue Impact Scale	e (FIS) and Short Form-36 (SF-36) scores at week					
	48 and last study						
	Definitions used	for endpoints: <u>Relapse</u> was defined as new or					
		l signs or symptoms lasting at least 24 h without					
		efined relapse constituted an increase of either 1					
		wo EDSS functional system scores, or 2 points in onal system score (excluding bowel and bladder					
		ebral function), or $0.5$ points in total EDSS score					
	from a previous of	clinically stable assessment time to 12 week sus-					
	-	ion of disability, defined as an increase from base-					
	line of at least 1 E	DSS point (or $\ge 0.5$ points when baseline EDSS					
	_	points that persisted for at least 12 weeks					
Follow-up		on in TOWER was variable and ended 48 weeks					
		ent was randomized into the study					
Treatment history	Mixed (based on e	exclusion criteria)					
Risk of bias							
Random sequence gen		Adequate.					
Allocation concealment	nt	Adequate "After a screening phase (up to 4					
	weeks), investigators used the allocation se-						
	quence to randomly assign eligible patients"						
Blinding of participan		Adequate.					
Blinding of outcome as	Adequate						
Incomplete outcome d		Adequate					
Intention to treat analysis							
Selective reporting None detected							
Other sources of bias		Funding: manufacturer					
runung: manufacturer							

**TENERE-((TErifluNomidE and REbifR) )study, Vermersch et al. 2014** (66), not included in Khai et al. (27)

<b>RCT identification</b>	NCT00883337							
Study setting		Rater-blinded study, randomized multicentre study						
Participants	<u>Eligibility criteria</u> : 18 years of age and older who met McDonald criteria for MS,13 had a relapsing clinical course with or without progression, and an Expanded Disability Status Scale (EDSS) score ≤5.5 at screening.14 Patients had to be relapse free for 30 days prior to randomisation. <u>Key exclusion criteria</u> : several restriction in previous and concom- itant medications, and relevant illnesses. <u>Baseline characteristics (group)</u> : Age 37+/-11; 68% female: EDSS							
	2,0+/-1,2							
Intervention group		14 mg oral once daily (n=111) 7 mg oral once daily (n=109)						
Comparison group	Interferon beta	n-1a 44mcg s.c three times/week (n=104)						
Outcome	of confirmed r any cause. See Scale (FIS) and tion (TSQM). <u>Definition used</u> or clinical wors ble for at least fever. required crease in at least	primary endpoint: time to failure, defined as first occurrence onfirmed relapse or permanent treatment discontinuation for cause. Secondary endpoints included ARR, Fatigue Impact e (FIS) and Treatment Satisfaction Questionnaire for Medica- (TSQM). <u>nition used for: <i>Relapse</i></u> criteria a new clinical sign/symptom inical worsening of a previous sign/symptom (previously sta- tor at least 30 days) that persisted for at least 24 hours without c. required a 1-point increase in each of two FS, a 2-point in- se in at least one FS (excluding bowel/bladder and cerebral) or increase of 0.5 points in EDSS score from the previous stable						
Follow-up	48 weeks after able duration of	the last patient was randomised, resulting in a vari-						
Treatment history		on exclusion criteria)						
Risk of bias	initia (bubbu c							
Random sequence gen	eration	Unclear, not described						
Allocation concealment		Unclear						
Blinding of participan nel	t and person-							
Blinding of outcome a		Adequate						
Incomplete outcome d	ata	<ul><li>22.4% discontinued treatment due to AEs</li><li>3 patients in IFN did not receive study drug.</li><li>Efficacy analyses: intention-to-treat population,</li><li>The safety analysis included all randomized pa-</li></ul>						
Selective reporting		tients exposed to study medication.						
Selective reportingUnclearOther sources of biasAuthors declare conflict of interest in form o laboration, employment or other with one or eral of the pharmaceutical companies								

### Appendix 3: Excluded studies and reasons for exclusions

#### **Information on the following tables:** CIS= Clinical Isolated Syndrome

CIS= Clinical Isolated Syndrome P= population I=Intervention C=Comparator S=Study design Y=Yes (the study fits that criteria) N=No (the study does not fit that criteria)

	CIS	Publica- tion date	Р	Ι	C	0	S	Exclu- sion/com- ments
Corrections to Safety and efficacy of fingolimod in pa- tients with relapsing-remitting multiple sclerosis		uuro					N	Exclude
(FREEDOMS II): A double-blind, randomised, pla- cebo-controlled, phase 3 trial. [Lancet Neurol 13 (2014) 545-56]. The Lancet Neurology 2014;13(6):536.								Correction up- dated in online version
Agius M, Meng X, Chin P, Grinspan A, Hashmonay R. Fingolimod therapy in early multiple sclerosis: An ef- ficacy analysis of the transforms and freedoms studies by time since first symptom. CNS Neuroscience and Therapeutics 2014;20(5):446-451.			N	Y	Y	Y	Y	Exclude subgroups of pa- tients <3 yrs since their first MS symptom
Arnold DL, Calabresi PA, Kieseier BC, Sheikh SI, Deykin A, Liu S, et al. Effect of peg-interferon beta-1a on MRI measures and freedom from measured dis- ease activity: 2-year results from the phase 3 AD- VANCE study. Mult Scler 2014;1):97.							N	Exclude Abstract
Arnold DL, Calabresi PA, Kieseier BC, Sheikh SI, Deykin A, Zhu Y, Liu S, You X, Sperling B, Hung S. Ef- fect of peg-interferon beta-1a on MRI measures and achieving no evidence of disease activity: results from a randomized controlled trial in relapsing-remitting multiple sclerosis. BMC Neurol. 2014 Dec 31;14(1):1058.			Y	Y	Y	N		Exclude ADVANCE Combined out- come of relapse and disability progression
Brinar V, Arnold DL, Cohen J, Coles AJ, Fox EJ, Hartung HP, et al. Alemtuzumab improves expanded disability status scale (EDSS) via effects on functional systems: CARE-MS II. Mult Scler 2013;1):283-284.							N	Exclude Abstract
Calabresi PA, Kieseier BC, Arnold DL, Balcer L, Boyko A, Pelletier J, et al. Clinical efficacy of peg-interferon beta-1a in relapsingremitting multiple sclerosis: 2- year data from the phase 3 ADVANCE study. Mult Scler 2014;1):42-43.							N	Exclude Abstract
Cascione M, Gaines C, Fang J, Dangond F, Miller A. Early and consistent reduction in relapses among pa- tients with relapsing-remitting multiple sclerosis re- ceiving subcutaneous interferon beta-1a: A post-hoc analysis of prisms data. Neurology 2014;1).							N	Exclude Abstract
Cascione M, Wynn D, Barbato LM, Pestreich L, Schofield L, McCague K. Randomized, open-label study to evaluate patient-reported outcomes with fin- golimod after changing from prior disease-modifying therapy for relapsing multiple sclerosis: EPOC study rationale and design. J Med Econ 2013;16(7):859- 865.						N		Exclude The comparator is disease-modi- fying therapies.

	CIS	Publica- tion	Р	Ι	C	0	S	Exclu- sion/com-
		date						ments
Chan A, Phillips JT, Fox RJ, Zhang A, Okwuokenye M,							Ν	Exclude
Kurukulasuriya NC. Differential recovery from relapse								
between treatment groups in the CONFIRM study of								Abstract
delayed-release dimethyl fumarate. Mult Scler 2014;1):110.								
Cofield SS, Gustafson T, Cutter GR, Wolinsky JS, Lu-	1						Ν	Exclude
blin FD. Physician and participant treatment guesses								
in the double-blind CombiRx study. Mult Scler								Abstract
2014;1):111-112.							<b>N</b> T	
Cohen JA, Belova A, Selmaj K, Wolf C, Oberye JJL, Van Den Tweel ERW, et al. Generic glatiramer acetate							Ν	Exclude
is equivalent to copaxone on efficacy and safety: Re-								Abstract
sults of the randomized doubleblind GATE trial in								
multiple sclerosis. Mult Scler 2014;1):38-39.								
Comi G, Freedman MS, Kappos L, Miller AE, Olsson							Ν	Exclude
TP, Wolinsky JS, et al. Effect of teriflunomide on lym-								A1
phocyte and neutrophil counts: Pooled analyses from four placebo-controlled studies. Mult Scler								Abstract
2014;1):93-94.								
Comi G, Martinelli V, Rodegher M, Moiola L, Leocani	Y							Exclude
L, Bajenaru O, et al. Effects of early treatment with								
glatiramer acetate in patients with clinically isolated								Not RRMS pa-
syndrome. Mult Scler 2013;19(8):1074-1083.							NT	tients
Comi G, Miller AE, Wolinsky JS, Benamor M, Bauer D, Truffinet P, et al. The effect of teriflunomide on							Ν	Exclude
lymphocyte and neutrophil count in patients with a								Abstract
first clinical episode consistent with multiple sclero-								instruct
sis: Results from the TOPIC study. J Neurol								
2014;261:S91.	ļ							
Confavreux C, Olsson TP, Comi G, Freedman MS, Mil- ler A, Wolinsky JS, et al. Teriflunomide hepatic safety							Ν	Exclude
results: Pooled data from three placebo-controlled								Abstract
studies. J Neurol 2013;260:S122.								instruct
Cutter G, Wolinsky JS, Comi G, Ladkani D, Knappertz							Ν	Exclude
V, Vainstein A, et al. Comparable clinical and MRI ef-								
ficacy of glatiramer acetate 40mg/mL TIW and								Abstract
20mg/mL QD: Results of a systematic review and meta-analysis. Mult Scler 2014;1):90-91.								
De Stefano N, Kappos L, Radue EW, Sprenger T, Piani							Ν	Exclude
Meier D, Haring D, et al. Fingolimod effect on diffuse								
tissue damage is partly independent of its effect on fo-								Abstract
cal damage in relapsingremitting multiple sclerosis								
patients. Mult Scler 2014;1):379. De Stefano N, Sprenger T, Freedman MS, Cree B, Sor-							N	Exclude
mani MP, Haring DA, et al. Including threshold rates							Ν	Exclude
of brain volume loss in the definition of disease-activ-								Abstract
ity-free in multiple sclerosis using fingolimod phase 3								
data. Mult Scler 2014;1):196-197.								
Deykin A, Arnold D, Hung S, Sheikh S, Seddighzadeh							Ν	Exclude
A, Zhu Y, et al. Interim analysis of 2-year clinical effi- cacy and safety of peg-interferon beta-1a in patients								Abstract
with relapsing-remitting multiple sclerosis: Data from								Abstract
the pivotal phase 3 advance study. Neurology 2014;1).								
Dhib-Jalbut S, Sumandeep S, Valenzuela R, Ito K, Pa-					Ν		Ν	Exclude
tel P, Rametta M. Immune response during interferon								Re-analysis of
beta-1b treatment in patients with multiple sclerosis								START study,
who experienced relapses and those who were re- lapse-free in the START study. J Neuroimmunol								which is obser- vational with
2013;254(1-2):131-140.								only interfreron
								(Betaseron)

	CIS	Publica- tion	Р	Ι	C	0	S	Exclu- sion/com-
		date						ments
Edan G, Kappos L, Montalban X, Polman C, Freed-		unte					N	Exclude
man M, Hartung H. Long term impact of early initia-								
tion of interferon beta-1B after a first clinical event								Abstract
suggestive of multiple sclerosis: Additional relapse								
rate, edss, and msss analyses after 8 years. 2013;80.								
Fox E, Edwards K, Burch JG, Kim E, Pestreich L,							Ν	Exclude
McCague K, et al. Treatment satisfaction and clinical								
improvement after switch to fingolimod. J Neurol								Abstract
2013;260:S126.							NT	<b>F</b>
Freedman M, Wolinsky J, Comi G, Kappos L, Olsson T, Miller A, et al. Long-term safety and efficacy of teri-							Ν	Exclude
flunomide in patients with relapsing forms of multiple								Abstract
sclerosis in the TEMSO extension trial. Mult Scler								Abstract
2013;1):225.								
Freedman M, Wolinsky J, Comi G, Kappos L, Olsson							Ν	Exclude
T, Miller A, et al. Safety and efficacy of teriflunomide								
for up to 9 years in relapsing forms of multiple sclero-								Abstract
sis: Update of the temso extension trial. Neurology								
2014;1).								
Freedman MS. Evidence for the efficacy of interferon	Y		Ν				Ν	Exclude
beta-1b in delaying the onset of clinically definite mul-								D
tiple sclerosis in individuals with clinically isolated								Review not SR
syndrome. Ther Adv Neurol Disord 2014;7(6):279- 288.								
Freedman MS, Ben-Amor AF, Issard D, Casset-Sema-							N	Exclude
naz F. Assessing a tool to predict disease activity in							IN	Exclude
patients with multiple sclerosis: A post-hoc analysis of								Abstract
clinical trial data on patients treated with subcutane-								isotiuot
ous interferon beta-1a. Mult Scler 2013;1):262.								
Freedman MS, Stefano N, Barkhof F, Polman CH,	Y							Exclude
Comi G, Uitdehaag BMJ, et al. Patient subgroup anal-								
yses of the treatment effect of subcutaneous inter-								Not RRMS pa-
feron beta-1a on development of multiple sclerosis in								tients
the randomized controlled REFLEX study. J Neurol								
2014;261(3):490-499. Havrdova E, Gold R, Fox R, Kappos L, Phillips JT,							N	Exclude
Zhang A. BG-12 (dimethyl fumarate) treatment for re-							IN	Exclude
lapsing-remitting multiple sclerosis (RRMS) increases								Abstract
the proportion of patients free of measured clinical								Abstract
and neuroradiologic disease activity in the phase 3								
studies. 2013;80.								
Hung S, Kieseier BC, Arnold DL, Balcer L, Boyko A,				1	1	1	Ν	Exclude
Pelletier J, et al. Peg-interferon beta-1a provides im-								
provements in clinical and radiological disease activ-								Abstract
ity in relapsing-remitting multiple sclerosis: Year 1								
findings from the phase 3 advance study. Mult Scler								
2014;20 (7):926.							N	Ell-
Hunter SF, Hunter HM, Kantor D. Phase 1 trial moni- toring response to alemtuzumab (ALE) in naive and					1		Ν	Exclude
ALE-experienced subjects with refractory multiple								Abstract
sclerosis (MS). Mult Scler 2013;1):265-266.								110011001
Hutchinson M, Bar-Or A, Fox RJ, Gold R, Giovannoni				1	1	1	N	Exclude
G, Kita M, et al. Effect of BG-12 (dimethyl fumarate)					1			
in subgroups of patients with relapsing-remitting					1			Abstract
multiple sclerosis: Findings from Two Phase 3 Studies					1			
(DEFINE and CONFIRM). Mult Scler 2013;19								
(5):682-683.								

	CIS	Publica- tion date	Р	I	C	0	S	Exclu- sion/com- ments
Hutchinson M, Fox RJ, Havrdova E, Kurukulasuriya		uate					Ν	Exclude
NC, Sarda SP, Agarwal S, et al. Efficacy and safety of								
BG-12 (dimethyl fumarate) and other disease-modify-								Systematic re-
ing therapies for the treatment of relapsing-remitting								view. Date of
multiple sclerosis: A systematic review and mixed								search
treatment comparison. Curr Med Res Opin 2014;30(4):613-627.								15/11/2012
Hutchinson M, Fox RJ, Phillips JT, Miller DH, Havr-							Ν	Exclude
dova E, Kita M, et al. Efficacy and safety of BG-12 (di-								
methyl fumarate) in relapsing-remitting multiple scle-								Abstract
rosis in the phase 3 CONFIRM study. Mult Scler								
2013;19 (5):683.								N D.C.
Kappos L, Cohen J, Collins W, De Vera A, Zhang-Au-							Ν	Not RCT
berson L, Ritter S, et al. Fingolimod in relapsing mul- tiple sclerosis: An integrated analysis of safety find-								
ings. Multiple sclerosis and Related Disorders								
2014;3(4):494-504.								
Kappos L, O'Connor PW, Polman CH, Vermersch P,							Ν	Not RCT
Wiendl H, Pace A, et al. Clinical effects of natalizumab								
on multiple sclerosis appear early in treatment course.								
J Neurol 2013;260(5):1388-1395.						NT		
Kaufman M, Cree BA, De Seze J, Fox RJ, Gold R, Hartung HP, et al. Radiologic MS disease activity dur-						Ν		Exclude Re-analysis of
ing natalizumab treatment interruption: findings from								RESTORE study
RESTORE. J Neurol 2015;262(2):326-336.								and others pla-
								cebo groups
Khan O, Rieckmann P, Boyko A, Selmaj K, Zivadinov							Ν	Exclude
R. A multinational, multicenter, randomized, placebo-								
controlled, double-blind study to assess the efficacy,								Abstract
safety, and tolerability of glatiramer acetate 40 mg in- jection three times a week in subjects with RRMS: Ef-								
ficacy and safety results of the gala study. Neurology								
2013;80 (1 MeetingAbstracts).								
Kita M, Fox R, Phillips JT, Arnold D, Bar-Or A, Yang							Ν	Exclude
M. Clinical and neuroradiologic efficacy of BG-12 (di-								
methyl fumarate) in us patients with relapsing-remit-								Abstract
ting multiple sclerosis (RRMS): An integrated analysis								
of the phase 3 DEFINE and confirm studies. 2013;80. Leist T, Freedman M, Benamor M, Truffinet P, Du-							N	Exclude
kovic D, Comi G. Pooled safety data from four pla-							IN	Exclude
cebo-controlled teriflunomide studies. Neurology								Abstract
2014;1).								
Leist T, Freedman M, Kappos L, Olsson T, Miller A,							Ν	Exclude
Wolinsky J, et al. Pooled safety data from three pla-								
cebo-controlled teriflunomide studies. Mult Scler								Abstract
2013;1):274-275. Leist TP, Freedman MS, Kappos L, Olsson TP, Miller							N	Exclude
AE, Wolinsky JS, et al. Three placebo-controlled teri-							14	Exclude
flunomide studies: Pooled safety data. Mult Scler								Abstract
2014;20 (7):933-934.								
Leist TP, Freedman MS, Kappos L, Olsson TP, Miller							Ν	Exclude
AE, Wolinsky JS, et al. Pooled safety analyses from						1		
the teriflunomide clinical development program. Mult						1		Abstract
Scler 2014;1):110-111. Lublin F, Cofield S, Cutter G, Salter A, Wang J, Con-							N	Exclude
wit R, et al. Edss changes in combirx: Blinded, 7-year						1	11	EACIUUC
extension results for progression and improvement.								Abstract
Neurology 2013;80 (1 MeetingAbstracts).								
Lublin F, Cofield S, Cutter G, Salter A, Wang J, Con-							Ν	Exclude
wit R, et al. Relapse activity in the combirx trial:								A1
Blinded, 7-year extension results. Neurology 2013;80 (1 MeetingAbstracts).								Abstract
(1 meetiligAustracis).	l		L	I	<u> </u>	I	<u> </u>	

	CIS	Publica-	Р	Ι	С	0	S	Exclu-	
		tion date						sion/com- ments	
Macdonell R, Lublin F, Comi G, Freedman MS, Kap-							Ν	Exclude	
pos L, Maurer M, et al. Teriflunomide reduces re- lapse-related sequelae, severe relapses, hospitalisa-								Abstract	
tions and corticosteroid use: Pooled data from the								hostiuet	
phase 3 TEMSO and TOWER studies. Mult Scler									
2013;1):512-513.			NT				N	Ell-	
Mantia LL, Vacchi L, Rovaris M, Di Pietrantonj C, Ebers G, Fredrikson S, et al. Interferon beta for sec-			Ν				Ν	Exclude Review of	
ondary progressive multiple sclerosis: a systematic re-								Secondary pro-	
view. J Neurol Neurosurg Psychiatry 2013;84(4):420-								gressive	
426 Maurer M, Van Wijmeersch B, De Seze J, Meca-Lal-							N	Exclude	
lana J, Bozzi S, Vermersch P. Significant and mean-							IN	Exclude	
ingful improvement in treatment satisfaction with								Abstract	
teriflunomide versus subcutaneous IFNB-1A in pa-									
tients with relapsing ms results from Tenere. Value									
Health 2014;17 (7):A403. Mikol D, Freedman MS, Goldman MD, Hartung HP,							N	Exclude	
Havrdova E, Jeffery D, et al. Correlations between pa-							11	Exclude	
tient-reported ambulatory function (MSWS-12) and								Abstract	
objective disability measurements in SPMS: Analysis									
of ASCEND baseline data. Mult Scler 2014;1):408. Mikol D, Freedman MS, Goldman MD, Hartung HP,				-		-	N	Exclude	
Havrdova E, Jeffery D, et al. Ascend study of natali-							IN	Exclude	
zumab efficacy on disability in patients with second-								Abstract	
ary progressive multiple sclerosis (SPMS): Baseline									
demographics and disease characteristics. Ann Neurol 2013;74:S59-S60.									
Mikol D, Freedman MS, Goldman MD, Hartung HP,							Ν	Exclude	
Havrdova E, Jeffery D, et al. ASCEND study of natali-								Literaue	
zumab efficacy on reducing disability in patients with								Abstract	
secondary progressive multiple sclerosis: Baseline de- mographics and disease characteristics. Mult Scler									
2013;1):507-508.									
Miller A, Kappos L, Comi G, Confavreux C, Freedman							Ν	Exclude	
M, Olsson T. Teriflunomide efficacy and safety in pa-									
tients with relapsing multiple sclerosis: Results from tower, a second, pivotal, phase 3 placebo-controlled								Abstract	
study. 2013;80.									
Miller A, Wolinsky J, Kappos L, Comi G, Freedman M,							Ν	Exclude	
Olsson T, et al. Topic: Efficacy and safety of once-daily									
oral teriflunomide in patients with first clinical epi- sode consistent with multiple sclerosis. Neurology								Abstract	
2014;1).									
Miller A, Wolinsky J, Kappos L, Comi G, Freedman							Ν	Exclude	
MS, Olsson T, et al. TOPIC main outcomes: Efficacy									
and safety of once-daily oral teriflunomide in patients with clinically isolated syndrome. Mult Scler								Abstract	
2013;1):25-26.									
Miller AE, Wolinsky JS, Kappos L, Comi G, Freedman	Y							Exclude	
MS, Olsson TP, et al. Oral teriflunomide for patients								monto	
with a first clinical episode suggestive of multiple scle- rosis (TOPIC): A randomised, double-blind, placebo-								TOPIC	
controlled, phase 3 trial. The Lancet Neurology								Not RRMS pa-	
2014;13(10):977-986.								tients	
Montalban X, Barkhof F, Comi G, Hartung HP, Kap-							Ν	Exclude	
pos L, Khatri B, et al. Long term efficacy of fingolimod in patients with relapsing-remitting multiple sclerosis								Abstract	
previously treated with interferon b-1a or disease								ADSILACI	
modifying therapies: A post hoc analysis of the									
TRANSFORMS 4.5 year extension study. J Neurol									
2013;260:S124-S125.	1	I			1		I		

	CIS	Publica- tion	Р	Ι	С	0	S	Exclu-
		date						sion/com- ments
Moses H, Freedman M, Kappos L, Miller A, Olsson T,		uate					N	Exclude
Wolinsky J. Pre-DEFINEd subgroups analyses of							1	Exclude
tower, a placebo-controlled phase 3 trial of terifluno-								Abstract
mide in patients with relapsing multiple sclerosis.								
2013;80.								
Nabavi M, Abolfazli R, Beladimoghadam N, Shahriari							Ν	Exclude
S, Hatami-Sadabadi F, Shati M, et al. A randomized								41 1 1
double blind non-inferiority study of efficacy, safety and tolerability of actorif versus rebif in patients with								Abstract
relapsing remitting ms. Neuroepidemiology 2013;41								
(3-4):259.								
Nagtegaal GJA, Pohl C, Wattjes MP, Hulst HE, Freed-	Y		Ν			Ν		Exclude
man MS, Hartung HP, et al. Interferon beta-1b re-								
duces black holes in a randomised trial of clinically								Not RRMS pa-
isolated syndrome. Mult Scler 2014;20(2):234-242.								tients
O'Connor P, Lublin F, Wolinsky J, Comi G, Confa-							Ν	Exclude
vreux C, Freedman M. Teriflunomide reduces relapse-								
related sequelae, hospitalizations and corticosteroid								Abstract
use: A post-HOC analysis of the phase 3 tower study. 2013;80.								
Olsson T, Comi G, Freedman M, Miller A, Wolinsky J,							Ν	Exclude
Truffinet P, et al. Patients free of clinical ms activity in							1	Exclude
temso and tower: Pooled analyses of two phase 3 pla-								Abstract
cebo-controlled trials. Neurology 2014;1).								
Pakpoor J, Disanto G, Altmann DR, Pavitt S, Turner							Ν	Exclude
B, Calado-Marta M, et al. Is there an increased cancer								
risk in people with relapsing multiple sclerosis taking								Abstract
cladribine? Mult Scler 2014;1):455.							N	Ell-
Phillips JT, Fox RJ, Gold R, Havrdova E, Kappos L, Raghupathi K, et al. An integrated analysis of safety							Ν	Exclude
and tolerability of BG-12 (dimethyl fumarate) in pa-								Abstract
tients with relapsing-remitting multiple sclerosis from								ribbliact
phase 2 and 3 placebo-controlled studies. J Neurol								
2013;260:S75.								
Stefano N, Comi G, Kappos L, Freedman MS, Polman	Y		Ν					Exclude
CH, Uitdehaag BMJ, et al. Efficacy of subcutaneous								
interferon beta-1a on MRI outcomes in a randomised								Not RRMS pa-
controlled trial of patients with clinically isolated syn- dromes. J Neurol Neurosurg Psychiatry								tients
2014;85(6):647-653.								
Svenningsson A, Sundstrom P, Salzer J, Vagberg M.							N	Exclude
MS disease activity in RESTORE: a randomized 24-								
week natalizumab treatment interruption study. Neu-								
rology 2014;83(22):2099-2100.								
Tenenbaum N, Schofield L, Meng X, Kern R. The pre-							Ν	Exclude
ferms study: Evaluating real-world patient retention								Abatra at
on oral fingolimod compared with injectable disease modifying therapies in relapsing-remitting multiple								Abstract
sclerosis. Neurology 2014;1).								
Tolley K, Hutchinson M, Pachner A, Kinter ET, Sper-				1		1	Ν	Exclude
ling B, You X, et al. Systematic literature review and								
network meta-analysis of peg-interferon beta-1a and								Abstract
injectable therapies for relapsing-remitting multiple								
sclerosis. Mult Scler 2014;1):209.				<u> </u>				
Tunde C. [Natalizumab retreatment: effectiveness and					Ν			Exclude
long-term safety in multiple sclerosis in the STRATA								Everybody get
study]. Ideggyogyaszati Szemle 2014;67(7-8):277-279.								

	CIS	Publica- tion date	Р	Ι	C	0	S	Exclu- sion/com- ments
Twyman C, Montalban X, Arnold D, Cohen J, Coles A,		uate					N	Exclude
Confavreux C, et al. Relapse outcomes with								Linerade
alemtuzumab vs IFNB-1Å in active relapsing-remit-								Abstract
ting multiple sclerosis patients who experienced dis-								
ease activity while on prior therapy (CARE-MS II).								
Neurology 2013;80 (1 MeetingAbstracts).								
White JT, Kieseier BC, Newsome SD, Zhu Y, Cui Y,							Ν	Exclude
Seddighzadeh A, et al. Immunogenicity with peg-in-								
terferon beta-1a in patients with relapsing-remitting								Abstract
multiple sclerosis: 2-year data from the randomised phase 3, multicentre ADVANCE study in relapsing-re-								
mitting multiple sclerosis. J Neurol 2014;261:S234.								
Wolinsky JS, Narayana PA, Nelson F, Datta S, O'Con-						Ν		Exclude
nor P, Confavreux C, et al. Magnetic resonance imag-						14		Exclude
ing outcomes from a phase III trial of teriflunomide.								Not our out-
Mult Scler 2013;19(10):1310-1319.								come
Wolinsky JS, Truffinet P, Bauer D, Miller AE. Efficacy							Ν	Exclude
of teriflunomide in patients with early stage MS: Anal-								
ysis of the TOPIC study using 2010 McDonald diag-								Abstract
nostic criteria. Mult Scler 2014;1):109-110.								
Wolinsky JS, Borresen TE, Dietrich DW, Wynn D, Sidi				Ν				All patients used
Y, Steinerman JR, Knappertz V, Kolodny S; GLACIER								glatiramer ace-
Study Group. GLACIER: An open-label, randomized,								tate 20 mg some
multicenter study to assess the safety and tolerability of glatiramer acetate 40 mg three-times weekly ver-								of them switched to gla-
sus 20 mg daily in patients with relapsing-remitting								tiramer acetate
multiple sclerosis. Mult Scler Relat Disord. 2015								40 mg
Jul;4(4):370-6								40 mg
Zagmutt F, Carroll C. A network meta-analysis as-							Ν	Exclude
sessing the rate of adverse events and drop outs of al-								
ternative treatments for relapsing forms of multiple								Abstract
sclerosis. Neurology 2013;80 (1 MeetingAbstracts).								
Zagmutt FJ, Carroll CA. Mixed treatment compa rison							Ν	Exclude
of adverse events for BG-12, glatiramer, and terifluno-								41
mide for the treatment of relapsing forms of multiple								Abstract
sclerosis. Value Health 2013;16 (7):A720. Zagmutt FJ, Carroll CA. Meta-analysis of adverse							N	Exclude
events in recent randomized clinical trials for dimethil							Ν	SR. Date of
fumarate, glatiramer acetate and teriflunomide for the								search:
treatment of relapsing forms of multiple sclerosis. Int								January 2013
J Neurosci 2014.								
Kieseier BC, Arnold DL, Balcer LJ, Boyko AA, Pelletier	l		Y	Y	Ν	Y	N	Exclude
J, Liu S, Zhu Y, Seddighzadeh A, Hung S, Deykin A,								
Sheikh SI, Calabresi PA. Peg-interferon beta-1a in								
multiple sclerosis: 2-year results from ADVANCE.								
Mult Scler. 2014								

### Appendix 4 Ongoing studies and other potential relevant literature

Below is the list of randomized control trials identifiend on the WHO ICTRP website. Due to the lack of information, we could not determine whether these studies fit our criteria of selection. These studies may add to the evidence.

1) Retinal Nerve Fiber Layer (RNFL) as measured by Optical Coherence Tomography (OCT) to Depict axonal loss in Early RRMS treated with difFEreNt dosage of subCutaneous IFN bEta 1a - DEFENCE

http://apps.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2009-015007-97-IT

3) Long-Term Safety and Efficacy Study of Oral BG00012 Monotherapy in Relapsing-Remitting Multiple Sclerosis

http://apps.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2008-004753-14-BE

4) ADVANCED MRI STUDY ON INFLAMMATORY AND DEGENRATIVE DAMAGE IN MULTIPLE SCLEROSIS - RMaIDSM http://apps.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2008-007162-32-IT

5) A Phase 3 Randomized, Rater- and Dose-Blinded Study Comparing Two Annual Cycles of Intravenous Low- and High-Dose Alemtuzumab to Three-Times Weekly Subcutaneous Interferon Beta-1a (Rebif®) in Patients with Relapsing-Remitting Multiple Scleroris Who Have Relapsed On Therapy - CARE MS-II

http://apps.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2007-001162-32-GB

6) Long-term extension of the multinational, double-blind, placebo controlled study EFC6049 (HMR1726D/3001) to document the safety of two doses of teriflunomide (7 and 14 mg) in patients with multiple sclerosis with relapses http://apps.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2006-003361-14-FI

7) A pilot multi-centre randomised controlled trial of sequential treatment with Mitoxantrone and Glatiramer Acetate vs. Interferon Beta-1a in early active relapsing remitting Multiple Sclerosis

http://apps.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2004-004903-39-GB 8) Study of Montelukast on Gastrointestinal Tolerability in Patients With Relapsing Forms of Multiple Sclerosis Receiving Tecfidera

http://apps.who.int/trialsearch/Trial2.aspx?TrialID=NCT02410278

9) Impact of Natalizumab versus Fingolimod on Central Nervous System (CNS) Tissue Damage and Recovery in Active Relapsing-Remitting Multiple Sclerosis (RRMS) Subjects

http://apps.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2013-004622-29-IT 10) A study to evaluate the effect of aspirin on flushing in patients with RRMS treated with Tecfidera http://apps.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2013-001895-40-IE

11) Study to investigate the ability of a blood-derived score to select patients with relapsing multiple sclerosis who benefit from treatment with human immune globulin http://apps.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2012-005086-12-AT

12) MS Study Evaluating Safety and Efficacy of Two Doses of Fingolimod Versus Copaxone

http://apps.who.int/trialsearch/Trial2.aspx?TrialID=NCT01633112

13) A Study of Ocrelizumab in Comparison With Interferon Beta-1a in Patients With Relapsing Multiple Sclerosis

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http://apps.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2010-020315-36-BE
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14) A 18-month, open-label, rater-blinded, randomized, multi-center, active-controlled, parallel-group pilot study to assess efficacy and safety of fingolimod (Gilenya) in comparison to interferon beta-1b in treating the cognitive symptoms associated to relapsing-remitting multiple sclerosis and to assess possible relationship of these effects to regional brain atrophy

http://apps.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2010-023023-19-IT

15) A Study of Ocrelizumab in Comparison With Interferon Beta-1a in Patients With Relapsing Multiple Sclerosis

http://apps.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2010-020337-99-GB

# **Appendix 5: GRADE evaluation of comparisons**

#### Interferon beta-1a 22 mcg compared to Placebo for RRMS

		Qu	ality assess	ment			Nº of pa	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Inter- feron beta-1a 22 mcg	Pla- cebo	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	lm- portance
Annuali	sed relaps	se rate										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	seri- ous <u>3</u>	none	-/189	-/187	<b>RR</b> 0.69 (0.57 to 0.83)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 123	
Disease	e Progress	ion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	very seri- ous 45	none	64/189 (33.9%)	77/187 (41.2%)	<b>RR</b> <b>0.84</b> (0.61 to 1.19)	66 fewer per 1000 (from 78 more to 161 fewer)	⊕⊕○○ LOW 1245	
Withdra	wal due to	adverse	e events									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	very seri- ous 45	none	6/189 (3.2%)	2/187 (1.1%)	<b>RR</b> <b>1.68</b> (0.50 to 5.98)	7 more per 1000 (from 5 fewer to 53 more)	⊕⊕○○ LOW 1245	

MD - mean difference, RR - relative risk

Only one study, not possible to check for inconsistency 1.

2 Patients were treatment naïve.

3. 4.

For continuous outcomes, a sample size of approximately 400 may not be sufficient to insure prognostic balance (ref GRADE 6 paper). The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).

5. Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper).

### Interferon beta-1a 30 mcg compared to Placebo for RRMS

		Qu	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Inter- feron beta-1a 30 mcg	Pla- cebo	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relaps	se rate										
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous 1	not se- rious	none	-/659	-/647	<b>RR</b> <b>0.76</b> (0.65 to 0.89)	0 fewer per 1000 (from 0 fewer to 0 fewer)		
Disease	e Progress	sion			•						•	
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous 1	seri- ous ²	none	70/605 (11.6%)	96/593 (16.2%)	<b>RR</b> <b>0.68</b> (0.50 to 0.95)	52 fewer per 1000 (from 8 fewer to 81 fewer)	MODER- ATE 12	
Withdra	wal due to	o adverse	e events		•	•		•				
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous 1	very seri- ous 23	none	34/659 (5.2%)	21/647 (3.2%)	<b>RR</b> <b>1.73</b> (0.82 to 3.87)	24 more per 1000 (from 6 fewer to 93 more)		

MD – mean difference, RR – relative risk

1.

2.

In the minor contributing study patients were treatment naïve. Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). 3.

#### Interferon beta-1a 44 mcg compared to Placebo for RRMS

		Qu	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Inter- feron beta-1a 44 mcg	Pla- cebo	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relap	se rate	•	-							•	
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous 1	not se- rious	none	-/204	-/247	<b>RR</b> 0.67 (0.54 to 0.80)	0 fewer per 1000 (from 0 fewer to 0 fewer)		
Disease	e Progress	sion	•	•								
1	ran- domis ed tri- als	not seri- ous	not seri- ous 2	not seri- ous ³	very seri- ous 45	none	54/184 (29.3%)	77/187 (41.2%)	<b>RR</b> 0.70 (0.48 to 1.04)	124 fewer per 1000 (from 16 more to 214 fewer)	⊕⊕○○ LOW 2345	
Withdra	awal due te	o advers	e events									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 2	not seri- ous ³	very seri- ous 46	none	9/184 (4.9%)	2/187 (1.1%)	<b>RR</b> 5.32 (1.09 to 41.63)	46 more per 1000 (from 1 more to 435 more)	⊕⊕○○ LOW 2346	

MD – mean difference, RR – relative risk

In the major contributing study patients were treatment naïve. Only one study, not possible to check for inconsistency 1.

2.

3. Patients were treatment naïve.

4.

Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). The effect estimate and the confidence interval are not robust. Minor variations in number of events would change the results. 5.

6.

#### Glatiramer acetate 20 mg compared to Placebo for RRMS

		Qu	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Glati- ramer acetate 20 mg	Pla- cebo	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relaps	se rate	•		•						•	
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous 1	not se- rious	none	-/595	-/609	<b>RR</b> <b>0.70</b> (0.60 to 0.82)	0 fewer per 1000 (from 0 fewer to 0 fewer)		
Disease	e Progress	ion	•		•						•	
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous ²	very seri- ous 34	none	83/475 (17.5%)	93/489 (19.0%)	<b>RR</b> <b>0.88</b> (0.61 to 1.21)	23 fewer per 1000 (from 40 more to 74 fewer)	⊕⊕○○ LOW 234	
Withdra	awal due to	adverse	e events									
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous 1	very seri- ous 34	none	43/595 (7.2%)	41/609 (6.7%)	<b>RR</b> <b>1.22</b> (0.64 to 2.66)	15 more per 1000 (from 24 fewer to 112 more)	⊕⊕○○ LOW 134	

MD – mean difference, RR – relative risk

1.

2.

3.

In the minor contributing studies, patients were treatment naïve or had an unclear treatment history. In the minor contributing study patients were treatment naïve. Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). 4.

### Glatiramer acetate 40 mg compared to Placebo for RRMS

		Qu	ality assess	ment			Nº of pa	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sidera- tions	Glati- ramer acetate 40 mg	Pla- cebo	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	lm- portance
Annualis	sed relaps	e rate		-								
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	not se- rious	none	-/943	-/461	<b>RR</b> 0.66 (0.52 to 0.82)	0 fewer per 1000 (from 0 fewer to 0 fewer)		
Withdra	wal due to	adverse	events									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous ²³	none	29/943 (3.1%)	6/461 (1.3%)	<b>RR</b> 2.50 (0.86 to 8.29)	20 more per 1000 (from 2 fewer to 95 more)		

MD - mean difference, RR - relative risk

Only one study, not possible to check for inconsistency
 Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper).
 The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).

### Dimethyl fumarate 240 mg two times daily compared to Placebo for RRMS

		Qua	ality assess	ment			Nº of pa	tients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rect ness	Impre- cision	Other con- sider- ation s	Dimethyl fumarate 240 mg two times daily	Pla- cebo	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relap	se rate										
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	not se- rious	none	-/769	-/771	<b>RR</b> <b>0.50</b> (0.42 to 0.60)	0 fewer per 1000 (from 0 fewer to 0 fewer)	⊕⊕⊕ _{HIGH}	
Disease	e Progress	sion										
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	not se- rious	none	113/768 (14.7%)	172/771 (22.3%)	<b>RR</b> <b>0.65</b> (0.49 to 0.85)	78 fewer per 1000 (from 33 fewer to 114 fewer)	⊕⊕⊕ _{HIGH}	
Withdra	awal due te	o advers	e events	I	,							
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	very seri- ous 12	none	109/769 (14.2%)	90/771 (11.7%)	<b>RR</b> 1.24 (0.74 to 2.13)	28 more per 1000 (from 30 fewer to 132 more)	⊕⊕OO LOW 12	

MD - mean difference, RR - relative risk

1.

Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). 2.

### Dimethyl fumarate 240 mg three times daily compared to Placebo for RRMS

		Qua	ality assess	ment			Nº of pa	tients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rect ness	lm- preci- sion	Other con- sider- ation s	Dimethyl fumarate 240 mg three times daily	Pla- cebo	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	lm- portance
Annuali	ised relap	se rate										
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	not seri- ous	none	-/761	-/771	<b>RR</b> <b>0.50</b> (0.42 to 0.60)	0 fewer per 1000 (from 0 fewer to 0 fewer)	⊕⊕⊕ _{HIGH}	
Disease	e Progress	sion										
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	not seri- ous	none	120/761 (15.8%)	172/771 (22.3%)	<b>RR</b> <b>0.68</b> (0.52 to 0.89)	71 fewer per 1000 (from 25 fewer to 107 fewer)	⊕⊕⊕ _{HIGH}	
Withdra	wal due to	o advers	e events									
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	very seri- ous 12	none	109/760 (14.3%)	93/771 (12.1%)	<b>RR</b> 1.25 (0.74 to 2.13)	30 more per 1000 (from 31 fewer to 136 more)		

MD - mean difference, RR - relative risk

1.

Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). 2.

### Teriflunomide oral 7 mg compared to Placebo for RRMS

		Qu	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Teri- fluno- mide oral 7 mg	Pla- cebo	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	lm- portance
Annuali	ised relaps	se rate										
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous 1	not se- rious	none	-/802	-/806	<b>RR</b> <b>0.73</b> (0.64 to 0.84)	0 fewer per 1000 (from 0 fewer to 0 fewer)		
Disease	e Progress	ion			•						•	
1	ran- domis ed tri- als	not seri- ous	not seri- ous ²	not seri- ous	very seri- ous 34	none	79/365 (21.6%)	99/363 (27.3%)	<b>RR</b> <b>0.80</b> (0.55 to 1.13)	55 fewer per 1000 (from 35 more to 123 fewer)	⊕⊕○○ LOW 234	
Withdra	wal due to	adverse	e events									
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous 1	very seri- ous 34	none	97/802 (12.1%)	57/806 (7.1%)	<b>RR</b> 1.54 (0.89 to 2.51)	38 more per 1000 (from 8 fewer to 107 more)	⊕⊕○○ LOW 134	

MD - mean difference, RR - relative risk

In the minor contributing study patients were treatment naïve.
 Only one study, not possible to check for inconsistency
 Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper).
 The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both no effect.

both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).

### Teriflunomide oral 14mg compared to Placebo for RRMS

		Qu	ality assess	ment			Nº of pa	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Teri- fluno- mide oral 14mg	Pla- cebo	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relaps	se rate										
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous 1	not se- rious	none	-/824	-/806	<b>RR</b> 0.67 (0.58 to 0.78)	0 fewer per 1000 (from 0 fewer to 0 fewer)		
Disease	e Progress	sion	•									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 2	not seri- ous	very seri- ous 34	none	72/358 (20.1%)	99/363 (27.3%)	<b>RR</b> <b>0.73</b> (0.51 to 1.05)	74 fewer per 1000 (from 14 more to 134 fewer)	⊕⊕○○ Low 234	
Withdra	wal due to	o advers	e events									
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous 1	very seri- ous 35	none	100/824 (12.1%)	57/806 (7.1%)	<b>RR</b> <b>1.70</b> (1.02 to 3.01)	50 more per 1000 (from 1 more to 142 more)	⊕⊕○○ LOW 135	

MD - mean difference, RR - relative risk

In the minor contributing study patients were treatment naïve.
 Only one study, not possible to check for inconsistency
 Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper).
 The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).
 The effect estimate and the confidence interval are not robust. Minor variations in number of events would change the results.

#### Fingolimod oral 0.5 mg compared to Placebo for RRMS

		Qu	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Fin- golimod oral 0.5 mg	Pla- cebo	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	sed relaps	se rate	•								•	<u>.</u>
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	not se- rious	none	-/840	-/830	<b>RR</b> <b>0.49</b> (0.41 to 0.57)	0 fewer per 1000 (from 0 fewer to 0 fewer)	⊕⊕⊕ _{HIGH}	
Disease	e Progress	sion	•								•	
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	not se- rious	none	124/783 (15.8%)	164/773 (21.2%)	<b>RR</b> 0.75 (0.56 to 0.98)	53 fewer per 1000 (from 4 fewer to 93 fewer)	⊕⊕⊕⊕ _{HIGH}	
Withdra	wal due to	o advers	e events				•				L	•
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	very seri- ous 12	none	104/840 (12.4%)	72/830 (8.7%)	<b>RR</b> 1.49 (0.86 to 2.50)	43 more per 1000 (from 12 fewer to 130 more)	LOW 12	

MD – mean difference, RR – relative risk

1.

Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). 2.

## Fingolimod oral 1.25 mg compared to Placebo for RRMS

		Qua	ality assessr	nent			Nº of pa	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	lm- pre- cisio n	Other con- sider- ations	Fin- golimod oral 1.25 mg	Placebo	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relaps	se rate	•					•				
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	not seri- ous	none	-/853	-/830	<b>RR</b> <b>0.43</b> (0.37 to 0.51)	0 fewer per 1000 (from 0 fewer to 0 fewer)	⊕⊕⊕ _{HIGH}	
Disease	e Progress	ion	•					•				
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	not seri- ous	none	119/799 (14.9%)	164/773 (21.2%)	<b>RR</b> <b>0.70</b> (0.52 to 0.92)	64 fewer per 1000 (from 17 fewer to 102 fewer)	⊕⊕⊕ _{HIGH}	
Withdra	wal due to	adverse	e events					•				
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	seri- ous 1	none	139/853 (16.3%)	72/830 (8.7%)	<b>RR</b> <b>1.93</b> (1.18 to 3.14)	81 more per 1000 (from 16 more to 186 more)		

MD – mean difference, RR – relative risk

1. Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper).

## Peg-interferon beta-1a 125 mcg once every two weeks compared to Placebo for RRMS

		Qua	ality assess	ment			Nº of pa	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rect ness	Impre- cision	Other con- sider- ations	Peg-in- terferon beta-1a 125 mcg once every two weeks	Pla- cebo	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relaps	se rate										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	not se- rious	none	-/512	-/500	<b>RR</b> <b>0.65</b> (0.49 to 0.85)	0 fewer per 1000 (from 0 fewer to 0 fewer)		
Disease	e Progress	sion					_					
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous ²³	none	31/512 (6.1%)	50/500 (10.0%)	<b>RR</b> <b>0.61</b> (0.36 to 0.98)	39 fewer per 1000 (from 2 fewer to 64 fewer)	⊕⊕○○ LOW 123	
Withdra	wal due to	o advers	e events									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous ²⁴	none	25/512 (4.9%)	7/500 (1.4%)	<b>RR</b> <b>3.57</b> (1.27 to 11.14)	36 more per 1000 (from 4 more to 142 more)	LOW 124	

MD - mean difference, RR - relative risk

Only one study, not possible to check for inconsistency Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). The effect estimate and the confidence interval are not robust. Minor variations in number of events would change the results. 1. 2.

- Control group risks or approximately or is as seen at the orbust. Minor variations in number of events would change the results. The effect estimate and the confidence interval are not robust. Minor variations in number of events would change the results. 3. 4.

#### Peg-interferon beta-1a 125 mcg once every four weeks compared to Placebo for RRMS

		Qua	ality assess	ment			Nº of pa	atients		Effect		
Nº of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rect ness	Impre- cision	Other con- sider- ations	Peg-in- terferon beta-1a 125 mcg once every four weeks	Pla- cebo	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relaps	se rate										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	not se- rious	none	-/500	-/500	<b>RR</b> <b>0.73</b> (0.56 to 0.95)	0 fewer per 1000 (from 0 fewer to 0 fewer)		
Disease	e Progress	sion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous 23	none	31/500 (6.2%)	50/500 (10.0%)	<b>RR</b> 0.62 (0.38 to 1.01)	38 fewer per 1000 (from 1 more to 62 fewer)		
Withdra	wal due to	o advers	e events									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous 24	none	24/500 (4.8%)	7/500 (1.4%)	<b>RR</b> 3.47 (1.25 to 10.90)	35 more per 1000 (from 4 more to 139 more)	⊕⊕○○ Low 124	

MD - mean difference, RR - relative risk

Only one study, not possible to check for inconsistency
 Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper).
 The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).
 The effect estimate and the confidence interval are not robust. Minor variations in number of events would change the results

#### Natalizumab 300 mg intravenous every four weeks compared to Placebo for RRMS

		Qu	ality assess	ment			Nº of pa	tients		Effect		
Nº of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Natali- zumab 300 mg intrave- nous every four weeks	Pla- cebo	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	sed relaps	se rate										
2	ran- domis ed tri- als	not seri- ous	serious 1	not seri- ous ²	not se- rious	none	-/673	-/358	<b>RR</b> 0.30 (0.25 to 0.36)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 12	
Disease	Progress	sion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 3	not seri- ous 4	seri- ous ⁵	none	107/627 (17.1%)	91/315 (28.9%)	<b>RR</b> <b>0.59</b> (0.42 to 0.84)	118 fewer per 1000 (from 46 fewer to 168 fewer)	MODER- ATE 345	
Withdra	wal due to	o advers	e events									
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous ²	very seri- ous 56	none	38/673 (5.6%)	15/358 (4.2%)	<b>RR</b> <b>1.22</b> (0.50 to 2.74)	9 more per 1000 (from 21 fewer to 73 more)	⊕⊕○○ Low 255	

MD - mean difference, RR - relative risk

Heterogeneity may be explained by differences in study setting. One study compared natalizumab with placebo over a two years period while the other tested treatment interruption in natalizumab users One study compared natalizumab with placebo over a two years period while the other tested treatment interruption in natalizumab users Only one study, not possible to check for inconsistency 1.

2.

3.

4. Patients' treatment history was unclear.

Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). 5.

6. The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).

## Interferon beta-1b 250 mcg SC every other day compared to Placebo for RRMS

		Qu	ality assess	ment			Nº of pa	tients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Inter- feron beta-1b 250 mcg SC every other day	Pla- cebo	Rela- tive (95% Cl)	Absolute (95% Cl)	Quality	lm- portance
Annuali	ised relaps	se rate										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	seri- ous <u>3</u>	none	-/124	-/122	<b>RR</b> 0.65 (0.51 to 0.83)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 123	
Disease	e Progress	sion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	very seri- ous 45	none	43/122 (35.2%)	56/122 (45.9%)	<b>RR</b> 0.77 (0.50 to 1.17)	106 fewer per 1000 (from 78 more to 230 fewer)	⊕⊕○○ LOW 1245	
Withdra	wal due to	o advers	e events		•		•					
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	very seri- ous 46	none	1/124 (0.8%)	10/122 (8.2%)	<b>RR</b> 0.070 (0.003 to 0.480)	76 fewer per 1000 (from 43 fewer to 82 fewer)	⊕⊕○○ LOW 1246	

MD - mean difference, RR - relative risk

Only one study, not possible to check for inconsistency 1.

2. Patients were treatment naïve.

Facents were treatment harve. For continuous outcomes, a sample size of approximately 400 may not be sufficient to insure prognostic balance (ref GRADE 6 paper). Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). The officient end the confidence interval exercises the term in the sum of the confidence interval (CI) includes both no effect. 3. 4.

5.

The effect estimate and the confidence interval are not robust. Minor variations in number of events would change the results. 6.

## Alemtuzumab 24 mg IV q.d compared to Alemtuzumab 12 mg IV q.d for RRMS

		Qu	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectne ss	Impre- cision	Other con- sider- ations	Alemtu- zumab 24 mg IV q.d	Alemtu- zumab 12 mg IV q.d	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	sed relaps	e rate										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	seri- ous ²³	seri- ous 4	none	-/110	-/112	<b>RR</b> <b>0.55</b> (0.35 to 0.86)	0 fewer per 1000 (from 0 fewer to 0 fewer)	⊕⊕○○ LOW 1234	
Disease	e Progress	ion (disa	bility sustain	ed for 6 m	onths)							
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	seri- ous 23	very seri- ous 56	none	10/110 (9.1%)	8/112 (7.1%)	<b>RR</b> <b>0.85</b> (0.40 to 1.65)	11 fewer per 1000 (from 43 fewer to 46 more)	⊕○○○ VERY LOW 12355	
Withdra	wal due to	adverse	events									
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not se- rious ⁷	very seri- ous 56	none	7/280 (2.5%)	16/539 (3.0%)	<b>RR</b> 0.88 (0.30 to 2.31)	4 fewer per 1000 (from 21 fewer to 39 more)		

MD - mean difference, RR - relative risk

Only one study, not possible to check for inconsistency 1.

2. Few patients could have received the intended three treatments's rounds. Alemtuzumab arms were suspended from 2005 as immune thrombocytopenic purpura developed in three patients, and one of them died (patients were recruited from 2002 to 2004). Patients were treatment naïve.

3

4

For continuous outcomes, a sample size of approximately 400 may not be sufficient to insure prognostic balance (ref GRADE 6 paper). Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and 5. control group risks of approximately 50% or greater (ref GRADE 6 paper). The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes

6. both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).

7. In the minor contributing study patients were treatment naïve. In the major contributing study patients were treatment experienced.

## Interferon beta-1a 44 mcg compared to Alemtuzumab 12 mg IV q.d for RRMS

		Qu	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectne ss	lm- pre- cisio n	Other con- sider- ations	Inter- feron beta-1a 44 mcg	Alemtu- zumab 12 mg IV q.d	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relaps	se rate									_	
3	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not se- ri- ous 23	not seri- ous	none	-/500	-/924	<b>RR</b> 2.22 (1.89 to 2.63)	0 fewer per 1000 (from 0 fewer to 0 fewer)	HIGH 123	
Disease	e Progress	sion										
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not se- ri- ous 23	seri- ous 4	none	113/529 (21.4%)	102/924 (11.0%)	<b>RR</b> 1.95 (1.45 to 2.59)	105 more per 1000 (from 50 more to 176 more)	MODER- ATE 234	
Withdra	wal due to	advers	e events	•								
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not se- ri- ous 23	seri- ous 4	none	39/500 (7.8%)	21/924 (2.3%)	<b>RR</b> <b>3.60</b> (1.88 to 7.34)	59 more per 1000 (from 20 more to 144 more)	MODER- ATE 234	

MD - mean difference, RR - relative risk

1. Some inconsistency. It might be explained by the fact that in one study alemtuzumab arms were suspended.

Included approximately the same proportion of treatment naïve and experienced patients. 2.

Included approximately the same proportion of treatment have and expendenced patients.
 In the minor contributing study, alemtuzumab arms were suspended from 2005 as immune thrombocytopenic purpura developed in three patients, and one of them died (Patients were recruited from 2002 to 2004).
 Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper).

## Interferon beta-1a 44 mcg compared to Alemtuzumab 24 mg IV q.d for RRMS

		Qu	ality assess	sment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectne ss	Impre- cision	Other con- sider- ations	Inter- feron beta-1a 44 mcg	Alem- tuzu- mab 24 mg IV q.d	Rela- tive (95% Cl)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relaps	se rate							·			
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	seri- ous ²³	seri- ous 4	none	-/111	-/110	<b>RR</b> <b>3.33</b> (1.94 to 5.79)	0 fewer per 1000 (from 0 fewer to 0 fewer)	⊕⊕○○ LOW 1234	
Disease	e Progress	sion		•								
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	seri- ous 23	very seri- ous 56	none	24/111 (21.6%)	10/110 (9.1%)	<b>RR</b> 2.15 (1.10 to 4.55)	105 more per 1000 (from 9 more to 323 more)	⊕     ○     ○     ○     ○     ○     ○     ○     ○     ○     ○     ○     ○     ○     ○     ○     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □     □    □    □    □    □□    □□    □□    □□    □□    □□    □	
Withdra	wal due to	o advers	e events									
2	ran- domis ed tri- als	not seri- ous	not seri- ous	seri- ous ⁷⁸	very seri- ous 56	none	28/313 (8.9%)	7/280 (2.5%)	<b>RR</b> <b>4.08</b> (1.69 to 11.42)	77 more per 1000 (from 17 more to 261 more)		

MD - mean difference, RR - relative risk

1

Only one study, not possible to check for inconsistency Few patients could have received the intended three treatments's rounds. Alemtuzumab arms were suspended from 2005 as immune thrombocyto-penic purpura developed in three patients, and one of them died (Patients were recruited from 2002 to 2004). Patients were treatment naïve. 2.

3.

For continuous outcomes, a sample size of approximately 400 may not be sufficient to insure prognostic balance (ref GRADE 6 paper). Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). 4. 5.

6. The effect estimate and the confidence interval are not robust. Minor variations in number of events would change the results.

In one of the two studies, few patients could have received the intended three treatments's rounds. Alemtuzumab arms were suspended from 2005 as immune thrombocytopenic purpura developed in three patients, and one of them died (Patients were recruited from 2002 to 2004). In the minor contributing study patients were treatment naïve. In the major contributing study patients were treatment experienced. 7.

8.

## Interferon beta-1a 44 mcg compared to Interferon beta-1a 22 mcg for RRMS

		Qu	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Inter- feron beta-1a 44 mcg	Inter- feron beta-1a 22 mcg	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	sed relaps	se rate				_					_	
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	seri- ous ³	none	-/184	-/189	<b>RR</b> <b>0.68</b> (0.56 to 0.83)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 123	
Disease	e Progress	iion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	very seri- ous 45	none	54/184 (29.3%)	64/189 (33.9%)	<b>RR</b> <b>0.92</b> (0.65 to 1.30)	27 fewer per 1000 (from 102 more to 119 fewer)	⊕⊕○○ LOW 1245	
Withdra	wal due to	advers	events									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	very seri- ous 45	none	9/184 (4.9%)	6/189 (3.2%)	<b>RR</b> <b>1.31</b> (0.40 to 4.36)	10 more per 1000 (from 19 fewer to 107 more)	⊕⊕○○ LOW 1245	

MD - mean difference, RR - relative risk

1. Only one study, not possible to check for inconsistency

2. Patients were treatment naïve.

Patients were treatment haive. For continuous outcomes, a sample size of approximately 400 may not be sufficient to insure prognostic balance (ref GRADE 6 paper). Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes 3. 4.

5. both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).

## Interferon beta-1a 44 mcg compared to Interferon beta-1a 30 mcg for RRMS

		Qua	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Inter- feron beta-1a 44 mcg	Inter- feron beta-1a 30 mcg	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relaps	se rate		_		_						
3	ran- domis ed tri- als	not seri- ous 1	not seri- ous	not seri- ous ²	not se- rious	none	-/424	-/423	<b>RR</b> 0.76 (0.63 to 0.93)	0 fewer per 1000 (from 0 fewer to 0 fewer)	$\bigoplus_{HIGH} \bigoplus_{12}$	
Disease	e Progress	sion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous ³	not seri- ous 4	very seri- ous 56	none	43/339 (12.7%)	49/338 (14.5%)	<b>RR</b> <b>0.89</b> (0.55 to 1.38)	16 fewer per 1000 (from 55 more to 65 fewer)	⊕⊕○○ Low 3456	
Withdra	wal due to	o adverse	events					•				
1	ran- domis ed tri- als	not seri- ous	not seri- ous ³	not seri- ous 4	very seri- ous 56	none	16/339 (4.7%)	14/337 (4.2%)	<b>RR</b> 1.15 (0.43 to 3.10)	6 more per 1000 (from 24 fewer to 87 more)	⊕⊕○○ LOW <u>3456</u>	

MD - mean difference, RR - relative risk

The major contributing study had no risk of bias issue. 1.

2. Patients' treatment history was unclear in all three studies

3.

Patients' treatment history was unclear in all three studies Only one study, not possible to check for inconsistency Patients' treatment history was unclear Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). 4. 5.

6.

#### Interferon beta-1a 60 mcg compared to Interferon beta-1a 30 mcg for RRMS

		Qu	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Inter- feron beta-1a 60 mcg	Inter- feron beta-1a 30 mcg	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	lm- portance
Annuali	sed relaps	se rate				_						
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	seri- ous ³	none	-/400	-/402	<b>RR</b> <b>1.05</b> (0.88 to 1.25)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 123	
Disease	Progress	ion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	very seri- ous 34	none	108/400 (27.0%)	109/402 (27.1%)	<b>RR</b> <b>0.99</b> (0.71 to 1.39)	3 fewer per 1000 (from 79 fewer to 106 more)	⊕⊕○○ LOW 1234	
Withdra	wal due to	adverse	e events									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	very seri- ous 34	none	64/400 (16.0%)	45/402 (11.2%)	<b>RR</b> <b>1.43</b> (0.66 to 3.11)	48 more per 1000 (from 38 fewer to 236 more)	⊕⊕○○ LOW 1234	

MD - mean difference, RR - relative risk

1. Only one study, not possible to check for inconsistency

2.

Patients' treatment history was unclear The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). 3.

4.

#### Glatiramer acetate 20 mg compared to Interferon beta-1a 30 mcg for RRMS

		Qua	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Glati- ramer acetate 20 mg	Inter- feron beta-1a 30 mcg	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	lm- portance
Annuali	sed relaps	se rate		÷								
2	ran- domis ed tri- als	not seri- ous 1	not seri- ous	not seri- ous ²	seri- ous ³	none	-/314	-/305	<b>RR</b> <b>0.79</b> (0.61 to 1.02)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 123	
Disease	Progress	ion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 4	not seri- ous <u>5</u>	very seri- ous 36	none	74/259 (28.6%)	61/250 (24.4%)	<b>RR</b> 1.18 (0.81 to 1.75)	44 more per 1000 (from 46 fewer to 183 more)	⊕⊕○○ LOW 3456	
Withdra	wal due to	adverse	events							·		
1	ran- domis ed tri- als	not seri- ous	not seri- ous 4	not seri- ous ⁵	very seri- ous 36	none	11/259 (4.2%)	17/250 (6.8%)	<b>RR</b> <b>0.61</b> (0.22 to 1.67)	27 fewer per 1000 (from 46 more to 53 fewer)	⊕⊕○○ LOW <u>3456</u>	

MD – mean difference, RR – relative risk

The major contributing study had no risk of bias issue 1.

Unclear treatment history in both studies. In the major contributing study patients were excluded if prior use of either interferon or glatiramer ace-2. tate.

tate. The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). Only one study, not possible to check for inconsistency Unclear treatment history, but patients were excluded if prior use of either interferon or glatiramer acetate. Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). 3.

4.

5. 6.

## Fingolimod oral 0.5 mg compared to Interferon beta-1a 30 mcg for RRMS

		Qu	ality assess	ment			Nº of pa	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Fin- golimod oral 0.5 mg	Inter- feron beta-1a 30 mcg	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	lm- portance
Annuali	sed relaps	se rate		-								
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	not se- rious	none	-/431	-/435	<b>RR</b> <b>0.48</b> (0.35 to 0.64)	0 fewer per 1000 (from 0 fewer to 0 fewer)		
Disease	Progress	ion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous ²³	none	27/431 (6.3%)	38/435 (8.7%)	<b>RR</b> <b>0.72</b> (0.42 to 1.17)	24 fewer per 1000 (from 15 more to 51 fewer)		
Withdra	wal due to	advers	e events									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous ²³	none	25/429 (5.8%)	34/431 (7.9%)	<b>RR</b> <b>1.28</b> (0.52 to 3.44)	22 more per 1000 (from 38 fewer to 192 more)	⊕⊕○○ Low 123	

MD - mean difference, RR - relative risk

Only one study, not possible to check for inconsistency
 Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper).
 The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).

## Fingolimod oral 1.25 mg compared to Interferon beta-1a 30 mcg for RRMS

		Qu	ality assess	ment			№ of pa	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Fin- golimod oral 1.25 mg	Inter- feron beta-1a 30 mcg	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	sed relaps	se rate		-								
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	not se- rious	none	-/426	-/435	<b>RR</b> <b>0.63</b> (0.46 to 0.90)	0 fewer per 1000 (from 0 fewer to 0 fewer)		
Disease	e Progress	sion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous ²³	none	34/426 (8.0%)	38/435 (8.7%)	<b>RR</b> <b>0.99</b> (0.58 to 1.60)	1 fewer per 1000 (from 37 fewer to 52 more)		
Withdra	wal due to	advers	e events	•								
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous 34	none	28/420 (6.7%)	34/431 (7.9%)	<b>RR</b> 2.44 (1.09 to 5.68)	114 more per 1000 (from 7 more to 369 more)	⊕⊕○○ LOW 134	

MD - mean difference, RR - relative risk

1.

2.

Only one study, not possible to check for inconsistency The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). The effect estimate and the confidence interval are not robust. Minor variations in number of events would change the results. 3.

4.

## Interferon beta-1b 250 mcg SC every other day compared to Interferon beta-1a 30 mcg for RRMS

		Qua	ality assess	ment			Nº of pa	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	lm- preci- sion	Other con- sider- ation s	Inter- feron beta-1b 250 mcg SC every other day	Inter- feron beta-1a 30 mcg	Rela- tive (95% Cl)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relap	se rate	•				•				<u>.</u>	
2	ran- domis ed tri- als	not seri- ous 1	not seri- ous	not seri- ous ²	seri- ous ³	none	-/126	-/126	<b>RR</b> <b>0.71</b> (0.53 to 0.91)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 123	
Disease	e Progress	sion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 4	not seri- ous 5	very seri- ous 67	none	13/96 (13.5%)	28/92 (30.4%)	<b>RR</b> 0.44 (0.23 to 0.82)	170 fewer per 1000 (from 55 fewer to 234 fewer)		
Withdra	wal due to	o advers	e events				•				<u>.</u>	
1	ran- domis ed tri- als	not seri- ous	not seri- ous 4	not seri- ous ⁵	very seri- ous 68	none	5/96 (5.2%)	1/92 (1.1%)	<b>RR</b> 6.27 (0.79 to 172.30)	57 more per 1000 (from 2 fewer to 1000 more)		

MD - mean difference, RR - relative risk

The major contributing study had no risk of bias issue 1.

2

In the major contributing study patients were treatment naïve. For continuous outcomes, a sample size of approximately 400 may not be sufficient to insure prognostic balance (ref GRADE 6 paper). Only one study, not possible to check for inconsistency Patients were treatment naïve. 3.

4.

5. 6. Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper).

The effect estimate and the confidence interval are not robust. Minor variations in number of events would change the results.

7. 8. The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).

#### Glatiramer acetate 20 mg compared to Interferon beta-1a 44 mcg for RRMS

		Qua	ality assess	ment			Nº of p	patients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Glati- ramer acetate 20 mg	Inter- feron beta-1a 44 mcg	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	lm- portance
Annuali	sed relaps	se rate			_	_						
2	ran- domis ed tri- als	not seri- ous 1	not seri- ous	not seri- ous ²	seri- ous ³	none	-/433	-/441	<b>RR</b> 1.02 (0.83 to 1.28)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 123	
Disease	e Progress	ion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 4	not seri- ous <u>5</u>	very seri- ous 36	none	33/378 (8.7%)	45/386 (11.7%)	<b>RR</b> <b>0.75</b> (0.46 to 1.21)	29 fewer per 1000 (from 24 more to 63 fewer)	⊕⊕○○ LOW <u>3456</u>	
Withdra	wal due to	adverse	events		•	•				·		
1	ran- domis ed tri- als	not seri- ous	not seri- ous 4	not seri- ous <u>5</u>	very seri- ous 36	none	19/378 (5.0%)	23/386 (6.0%)	<b>RR</b> <b>0.88</b> (0.36 to 1.94)	7 fewer per 1000 (from 38 fewer to 56 more)	⊕⊕○○ LOW <u>3456</u>	

MD - mean difference, RR - relative risk

The major contributing study had no risk of bias issue 1.

2.

In the major contributing study had no fact blas issue In the major contributing study patients were treatment naïve. Treatment history was unclear in the other The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). Only one study, not possible to check for inconsistency Patients were treatment naïve. 3.

4.

5.

Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). 6.

#### Teriflunomide 7 mg oral compared to Interferon beta-1a 44 mcg SC t.i.w. for RRMS

		Qu	ality assess	ment			Nº of	patients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Teri- fluno- mide 7 mg oral	Inter- feron beta-1a 44 mcg SC t.i.w.	Rela- tive (95% Cl)	Absolute (95% Cl)	Quality	lm- portance
Annuali	sed relaps	e rate										_
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	seri- ous ²	none	-/109	-/104	<b>RR</b> <b>1.72</b> (1.24 to 2.44)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 12	
Withdra	wal due to	adverse	e events									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous 34	none	9/110 (8.2%)	22/101 (21.8%)	<b>RR</b> <b>0.40</b> (0.14 to 1.00)	131 fewer per 1000 (from 0 fewer to 187 fewer)		

MD - mean difference, RR - relative risk

Only one study, not possible to check for inconsistency
 For continuous outcomes, a sample size of approximately 400 may not be sufficient to insure prognostic balance (ref GRADE 6 paper).
 Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper).
 The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes but the context of the context of

both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).

## Teriflunomide 14 mg oral compared to Interferon beta-1a 44 mcg SC t.i.w. for RRMS

		Qu	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Teri- fluno- mide 14 mg oral	Inter- feron beta-1a 44 mcg SC t.i.w.	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	lm- portance
Annuali	sed relaps	e rate										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous ²³	none	-/111	-/104	<b>RR</b> <b>0.91</b> (0.62 to 1.36)	0 fewer per 1000 (from 0 fewer to 0 fewer)		
Withdra	wal due to	adverse	e events									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous 34	none	12/110 (10.9%)	22/101 (21.8%)	<b>RR</b> <b>0.54</b> (0.20 to 1.38)	100 fewer per 1000 (from 83 more to 174 fewer)	⊕⊕○○ LOW <u>134</u>	

MD – mean difference, RR – relative risk

Only one study, not possible to check for inconsistency
 For continuous outcomes, a sample size of approximately 400 may not be sufficient to insure prognostic balance (ref GRADE 6 paper).
 The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).
 Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and

control group risks of approximately 50% or greater (ref GRADE 6 paper).

## Interferon beta-1b 250 mcg SC every other day compared to Interferon beta-1a 44 mcg SC t.i.w. for RRMS

		Qua	ality assess	ment			Nº of pa	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Inter- feron beta-1b 250 mcg SC every other day	Inter- feron beta- 1a 44 mcg SC t.i.w.	Rela- tive (95% Cl)	Absolute (95% Cl)	Quality	Im- portance
Annuali	sed relaps	e rate										
1	ran- domis ed tri- als	seri- ous 1	not seri- ous ²	not seri- ous ³	very seri- ous 45	none	-/30	-/30	<b>RR</b> <b>0.81</b> (0.46 to 1.43)	0 fewer per 1000 (from 0 fewer to 0 fewer)		

MD – mean difference, RR – relative risk

- Insufficient reporting for randomization, and differences in baseline characteristics between groups 1.
- 2. Only one study, not possible to check for inconsistency

3. 4. Patients' treatment history was unclear.

- For continuous outcomes, a sample size of approximately 400 may not be sufficient to insure prognostic balance (ref GRADE 6 paper). The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). 5.

## Dimethyl fumarate 240 mg two times daily compared to Glatiramer acetate 20 mg for RRMS

		Qua	ality assess	ment			Nº of pa	tients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rect ness	lm- preci- sion	Other con- sider- ation s	Dimethyl fumarate 240 mg two times daily	Glati- ramer acetate 20 mg	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	lm- portance
Annuali	ised relap	se rate										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	not seri- ous	none	-/359	-/351	<b>RR</b> <b>0.59</b> (0.38 to 0.90)	0 fewer per 1000 (from 0 fewer to 0 fewer)		
Disease	e Progress	sion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous ²³	none	47/359 (13.1%)	56/350 (16.0%)	<b>RR</b> <b>0.78</b> (0.52 to 1.18)	35 fewer per 1000 (from 29 more to 77 fewer)		
Withdra	awal due te	o advers	e events	•								
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous 23	none	44/359 (12.3%)	35/351 (10.0%)	<b>RR</b> 1.18 (0.49 to 2.84)	18 more per 1000 (from 51 fewer to 183 more)		

MD - mean difference, RR - relative risk

Only one study, not possible to check for inconsistency
 Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper).
 The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).

## Dimethyl fumarate 240 mg three times daily compared to Glatiramer acetate 20 mg for RRMS

		Qua	ality assess	ment			Nº of pa	tients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rect ness	lm- preci- sion	Other con- sider- ation s	Dimethyl fumarate 240 mg three times daily	Glati- ramer acetate 20 mg	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	lm- portance
Annuali	ised relap	se rate	•									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	not seri- ous	none	-/345	-/350	<b>RR</b> <b>0.53</b> (0.35 to 0.79)	0 fewer per 1000 (from 0 fewer to 0 fewer)		
Disease	e Progress	sion	•									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous ²³	none	45/345 (13.0%)	56/350 (16.0%)	<b>RR</b> <b>0.79</b> (0.53 to 1.16)	34 fewer per 1000 (from 26 more to 75 fewer)		
Withdra	awal due te	o advers	e events									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous 23	none	41/344 (11.9%)	35/351 (10.0%)	<b>RR</b> 1.15 (0.52 to 2.56)	15 more per 1000 (from 48 fewer to 156 more)	⊕⊕○○ Low 123	

MD – mean difference, RR – relative risk

1.

Only one study, not possible to check for inconsistency Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and 2. control group risks of approximately 50% or greater (ref GRADE 6 paper). The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes

3. both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).

## Interferon beta-1b 250 mcg SC every other day compared to Glatiramer acetate 20mg for RRMS

		Qua	ality assess	ment			Nº of pa	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Inter- feron beta-1b 250 mcg SC every other day	Glati- ramer acetate 20mg	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	sed relaps	se rate										
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous 1	seri- ous ²	none	-/933	-/487	<b>RR</b> <b>1.07</b> (0.90 to 1.27)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 12	
Disease	e Progress	ion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 3	not seri- ous 1	seri- ous ²	none	188/897 (21.0%)	90/448 (20.1%)	<b>RR</b> <b>1.04</b> (0.74 to 1.46)	8 more per 1000 (from 52 fewer to 92 more)	MODER- ATE 123	
Withdra	wal due to	adverse	e events									
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous <u>1</u>	very seri- ous ²⁴	none	17/933 (1.8%)	12/487 (2.5%)	<b>RR</b> <b>0.91</b> (0.37 to 2.27)	2 fewer per 1000 (from 16 fewer to 31 more)	LOW 124	

MD – mean difference, RR – relative risk

Patients were treatment naïve. 1.

2. The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes

The control of a province of GRADE 6 paper).
 Only one study, not possible to check for inconsistency
 Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper).

## Interferon beta-1b 500 mcg SC every other day compared to Glatiramer acetate 20mg for RRMS

		Qu	ality assess	ment			№ of pa	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Inter- feron beta-1b 500 mcg SC every other day	Glati- ramer acetate 20mg	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	sed relaps	se rate										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	seri- ous <u>3</u>	none	-/899	-/448	<b>RR</b> 0.95 (0.80 to 1.12)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 123	
Disease	e Progress	ion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	seri- ous <u>3</u>	none	198/899 (22.0%)	90/448 (20.1%)	<b>RR</b> <b>1.01</b> (0.74 to 1.36)	2 more per 1000 (from 52 fewer to 72 more)	MODER- ATE 123	
Withdra	wal due to	advers	e events			•						
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	very seri- ous 34	none	20/899 (2.2%)	8/448 (1.8%)	<b>RR</b> <b>1.16</b> (0.46 to 3.05)	3 more per 1000 (from 10 fewer to 37 more)	⊕⊕○○ LOW 1234	

MD – mean difference, RR – relative risk

Only one study, not possible to check for inconsistency Patients were treatment naïve. 1.

2.

The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). 3.

Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper).

## Dimethyl fumarate 240 mg three times daily compared to Dimethyl fumarate 240 mg two times daily for RRMS

		Qua	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rect ness	lm- preci- sion	Other con- sider- ation s	Dimethyl fumarate 240 mg three times daily	Dimethyl fumarate 240 mg two times daily	Rel- ative (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relap	se rate										
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	seri- ous <u>1</u>	none	-/760	-/769	<b>RR</b> <b>1.01</b> (0.82 to 1.23)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 1	
Disease	e Progres	sion										
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	very seri- ous 12	none	120/761 (15.8%)	113/768 (14.7%)	<b>RR</b> 1.06 (0.78 to 1.42)	9 more per 1000 (from 32 fewer to 62 more)	LOW 12	
Withdra	wal due t	o advers	e events									
2	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	very seri- ous 12	none	109/760 (14.3%)	109/769 (14.2%)	<b>RR</b> <b>1.01</b> (0.58 to 1.73)	1 more per 1000 (from 60 fewer to 103 more)		

MD – mean difference, RR – relative risk

The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper). Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper). 1.

2.

## Teriflunomide oral 14 mg compared to Teriflunomide oral 7 mg for RRMS

		Qu	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Teri- fluno- mide oral 14 mg	Teri- fluno- mide oral 7 mg	Rela- tive (95% Cl)	Absolute (95% Cl)	Quality	Im- portance
Annuali	sed relaps	se rate										
4	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous <u>1</u>	seri- ous ²	none	-/935	-/912	<b>RR</b> 0.86 (0.74 to 1.00)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 12	
Disease	e Progress	ion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous ³	not seri- ous	very seri- ous ²⁴	none	72/358 (20.1%)	79/365 (21.6%)	<b>RR</b> <b>0.92</b> (0.64 to 1.35)	17 fewer per 1000 (from 76 more to 78 fewer)	⊕⊕○○ Low 234	
Withdra	wal due to	adverse	e events									
4	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous <u>1</u>	seri- ous ²⁴	none	112/934 (12.0%)	106/912 (11.6%)	<b>RR</b> 1.12 (0.73 to 1.85)	14 more per 1000 (from 31 fewer to 99 more)	MODER- ATE 124	

MD - mean difference, RR - relative risk

In the minor contributing study, patients were treatment naïve
 The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).
 Only one study, not possible to check for inconsistency
 Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and control group risks of approximately 50% or greater (ref GRADE 6 paper).

## Fingolimod oral 1.25 mg compared to Fingolomid oral 0.5 mg for RRMS

		Qua	ality assess	nent			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	lm- pre- cisio n	Other con- sider- ations	Fin- golimod oral 1.25 mg	Fingol- omid oral 0.5 mg	Rela- tive (95% Cl)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relaps	se rate	•	-		•	<u>.</u>		-	•		<u>.</u>
4	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	seri- ous ³	none	-/1273	-/1269	<b>RR</b> <b>0.98</b> (0.83 to 1.17)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 123	
Disease	e Progress	sion	•			•				•		
3	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous	seri- ous ³	none	153/1225 (12.5%)	151/1214 (12.4%)	<b>RR</b> <b>1.01</b> (0.78 to 1.32)	1 more per 1000 (from 27 fewer to 40 more)	MODER- ATE 3	
Withdra	wal due to	o advers	e events									•
4	ran- domis ed tri- als	not seri- ous	not seri- ous	not seri- ous ²	seri- ous ³	none	181/1273 (14.2%)	128/1269 (10.1%)	<b>RR</b> <b>1.43</b> (0.94 to 2.21)	43 more per 1000 (from 6 fewer to 122 more)	MODER- ATE 23	

MD – mean difference, RR – relative risk

1. 2. 3.

Some inconsistency. It may be explained by different definitions of relapse in studies In the minor contributing study, patients' treatment history was unclear. The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).

## Peg-interferon beta-1a 125 mcg once every four weeks compared to Peginterferon beta-1a 125 mcg once every two weeks for RRMS

		Qu	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Peg-in- terferon beta-1a 125 mcg once every four weeks	Peg-in- terferon beta-1a 125 mcg once every two weeks	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relaps	se rate	•		•		•					
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	seri- ous ²	none	-/500	-/512	<b>RR</b> <b>1.13</b> (0.84 to 1.52)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 12	
Disease	e Progress	sion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous 23	none	31/500 (6.2%)	31/512 (6.1%)	<b>RR</b> <b>1.02</b> (0.61 to 1.74)	1 more per 1000 (from 24 fewer to 45 more)		
Withdra	wal due to	o advers	e events									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous	very seri- ous 23	none	24/500 (4.8%)	25/512 (4.9%)	<b>RR</b> <b>0.98</b> (0.41 to 2.37)	1 fewer per 1000 (from 29 fewer to 67 more)	LOW 123	

MD - mean difference, RR - relative risk

Only one study, not possible to check for inconsistency
 The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper).
 Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and

control group risks of approximately 50% or greater (ref GRADE 6 paper).

## Interferon beta-1b 250 mcg SC every other day compared to Natalizumab 300 mg intravenous every 4 weeks for RRMS

		Qu	ality assess	ment			Nº of p	patients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Inter- feron beta-1b 250 mcg SC every other day	Natali- zumab 300 mg intrave- nous every 4 weeks	Rela- tive (95% CI)	Absolute (95% Cl)	Quality	Im- portance
Annuali	sed relaps	e rate							-			
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	seri- ous ²	very seri- ous 34	none	-/9	-/10	not esti- mabl e		VERY LOW 1234	

MD - mean difference, RR - relative risk

Only one study, not possible to check for inconsistency Study included only patients treated with natalizumab randomised to continue natalizumab or to switch to interferon. Patients selected into the studies may be different from the general MS population. 1. 2.

3. No meaningful information was given to be able to estimate the relative risk (the RR was 1.65*10^8(4510 to 2.52*10^9)

No meaningful information was given to be able to estimate the relative risk (the KK was 1.00 10 0(4010 to 2.02 10 0). For continuous outcomes, a sample size of approximately 400 may not be sufficient to insure prognostic balance (ref GRADE 6 paper). 4.

## Interferon beta-1b 500 mcg SC every other day compared to Interferon beta-1b 250 mcg SC every other day for RRMS

		Qu	ality assess	ment			Nº of p	atients		Effect		
№ of stud- ies	Study de- sign	Risk of bias	Incon- sistency	Indi- rectn ess	Impre- cision	Other con- sider- ations	Inter- feron beta-1b 500 mcg SC every other day	Inter- feron beta-1b 250 mcg SC every other day	Rela- tive (95% Cl)	Absolute (95% Cl)	Quality	Im- portance
Annuali	ised relaps	se rate										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	seri- ous ³	none	-/899	-/897	<b>RR</b> <b>0.93</b> (0.80 to 1.10)	0 fewer per 1000 (from 0 fewer to 0 fewer)	MODER- ATE 123	
Disease	e Progress	sion										
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	seri- ous ³	none	198/899 (22.0%)	188/897 (21.0%)	<b>RR</b> <b>1.10</b> (0.84 to 1.51)	21 more per 1000 (from 34 fewer to 107 more)	MODER- ATE 123	
Withdra	wal due to	o advers	e events									
1	ran- domis ed tri- als	not seri- ous	not seri- ous 1	not seri- ous ²	very seri- ous 34	none	20/899 (2.2%)	13/897 (1.4%)	<b>RR</b> <b>1.63</b> (0.66 to 4.11)	9 more per 1000 (from 5 fewer to 45 more)	⊕⊕○○ LOW 1234	

MD - mean difference, RR - relative risk

1. Only one study, not possible to check for inconsistency

Patients were treatment naive.
 The confidence interval (CI) includes both no effect, and the 25% threshold for appreciable benefit or harm. This corresponds to a CI that includes both 1.00, and 0.75 or 1.25 (ref GRADE 6 paper)
 Optimal information size (OIS) not met. For example, at least 200 events are required to meet OIS, based on a 25% relative risk reduction and

control group risks of approximately 50% or greater (ref GRADE 6 paper).

# Appendix 6: Full network meta-analysis results

# A6.1: Annualised relapse rate

Treatment	Placebo	Alemtuzumab 12 mg IV q.d	Alemtuzumab 24 mg IV q.d	Interferon beta-1a 22 mcg SC t.i.w	Interferon beta-1 a 3 0 m cg IM q.w			glatiramer acetate 20mg q.d	glatiramer acetate 40mg t.i.w			Teriflunomide or al 7 mg 14 mg	Fingolimod oral 0.5 mg	Fingolim od oral 1.25 mg	Peginterferon beta-1 a 125 mcg once every 2 weeks	1a 125 mcg once	Interferon beta- 1 b 250 mcg SC ev ery other day
Placebo		1								Ì							
Alemtuzumab 12 mg IV q.d	0.29 (0.23 to 0.35)	1															
Alemtuzumab 2.4 mg IV q.d	0.16 (0.1 to 0.25)	0.55 (0.35 to 0.86)	1	1													
Interferon beta-1a 22 mcg SC t.i.w	0.69 (0.57 to 0.83)	2.4 (1.9 to 3.12)	4.35 (2.71 to 7.13)		1												
Interferon beta-1a 30 mcg IM q.w	0.82 (0.73 to 0.91)	2.82 (2.33 to 3.59)	5.15 (3.24 to 8.29)	1.18 (0.97 to 1.46)													
Interferon beta-1a 44 m cg SC t.i.w	0.64 (0.56 to 0.72)	2.21 (1.9 to 2.64)	4.02 (2.6 to 6.34)	0.92 (0.76 to 1.11)	0.78 (0.68 to 0.89)												
Interferon beta-1a 60 m cg IM q.w	0.86 (0.7 to 1.06)	2.96 (2.31 to 4.02)	5.41 (3.3 to 8.99)	1.24 (0.96 to 1.63)	1.05 (0.88 to 1.25)	1.34 (1.09 to 1.7)	1										
glatiramer acetate 20mg q.d	0.65 (0.59 to 0.73)	2.25 (1.85 to 2.87)	4.1 (2.59 to 6.63)	0.94 (0.77 to 1.16)	0.8 (0.7 to 0.91)	1.02 (0.9 to 1.18)	0.76 (0.61 to 0.94)		1								
glatiramer acetate 40mg t.i.w	0.66 (0.52 to 0.82)	2.27 (1.7 to 3.14)	4.14 (2.47 to 6.95)	0.95 (0.72 to 1.27)	0.8 (0.62 to 1.02)	1.03 (0.8 to 1.33)	0.77 (0.56 to 1.03)	1.01 (0.78 to 1.28)		ı l							
dimethyl fumarate 240 mg two times daily	0.5 (0.42 to 0.6)	1.73 (1.35 to 2.31)	3.15 (1.92 to 5.23)	0.72 (0.56 to 0.93)	0.61 (0.49 to 0.75)	0.78 (0.63 to 0.97)	0.58 (0.44 to 0.76)	0.77 (0.63 to 0.93)	0.76 (0.57 to 1.01)	1							
dimethyl fumarate 240 mg three times daily	0.5 (0.42 to 0.6)	1.73 (1.35 to 2.33)	3.16 (1.93 to 5.28)	0.72 (0.57 to 0.94)	0.62 (0.5 to 0.75)	0.79 (0.64 to 0.98)	0.58 (0.45 to 0.76)	0.77 (0.64 to 0.93)	0.77 (0.58 to 1.02)	1.01 (0.82 to 1.23)	1	1					
Teriflunomide oral 7 mg	0.77 (0.68 to 0.9)	2.68 (2.13 to 3.53)	4.89 (3.03 to 7.93)	1.12 (0.9 to 1.42)	0.95 (0.8 to 1.13)	1.21 (1.02 to 1.47)	0.9 (0.71 to 1.16)	1.19 (1. to 1.42)	1.18 (0.91 to 1.55)	1.55 (1.24 to 1.96)	1.54 (1.23 to 1.94)	1					
Teriflunomide oral 14 mg	0.67 (0.58 to 0.77)	2.3 (1.83 to 3.03)	4.19 (2.6 to 6.9)	0.96 (0.77 to 1.22)	0.82 (0.68 to 0.98)	1.04 (0.87 to 1.27)	0.78 (0.6 to 0.99)	1.02 (0.85 to 1.22)	1.02 (0.78 to 1.33)	1.33 (1.06 to 1.68)	1.33 (1.05 to 1.67)	0.86 (0.74 to 1.)					
Fingolim od oral 0.5 mg	0.46 (0.39 to 0.54)	1.6 (1.25 to 2.09)	2.91 (1.79 to 4.79)	0.67 (0.53 to 0.85)	0.57 (0.47 to 0.67)	0.72 (0.6 to 0.88)	0.54 (0.42 to 0.68)	0.71 (0.59 to 0.85)	0.7 (0.54 to 0.92)	0.92 (0.73 to 1.17)	0.92 (0.73 to 1.16)	0.6 (0.48 to 0.73) 0.69 (0.56 to 0.85	1				
Fingolim od oral 1.25 mg	0.45 (0.39 to 0.53)	1.57 (1.23 to 2.06)	2.86 (1.76 to 4.66)	0.65 (0.52 to 0.83)	0.55 (0.47 to 0.66)	0.71 (0.58 to 0.87)	0.53 (0.41 to 0.67)	0.69 (0.57 to 0.83)	0.69 (0.53 to 0.9)	0.9 (0.71 to 1.15)	0.9 (0.71 to 1.14)	0.59 (0.47 to 0.71) 0.68 (0.55 to 0.84	0.98 (0.83 to 1.17)	1	ı l		
Peginterferon beta-1a 125 mcg once every 2 weeks	0.65 (0.49 to 0.85)	2.23 (1.6 to 3.19)	4.07 (2.33 to 7.07)	0.93 (0.67 to 1.29)	0.79 (0.58 to 1.06)	1.01 (0.74 to 1.36)	0.75 (0.53 to 1.05)	0.99 (0.73 to 1.32)	0.98 (0.69 to 1.41)	1.29 (0.93 to 1.8)	1.29 (0.92 to 1.77)	0.83 (0.6 to 1.13) 0.97 (0.7 to 1.32)	1.4 (1.02 to 1.92)	1.43 (1.03 to 1.95)	1		
peginterferon beta-1 a 125 mcg once every 4 weeks	0.73 (0.56 to 0.95)	2.52 (1.83 to 3.59)	4.59 (2.68 to 7.94)	1.06 (0.76 to 1.46)	0.89 (0.66 to 1.2)	1.14 (0.85 to 1.54)	0.85 (0.6 to 1.19)	1.12 (0.83 to 1.5)	1.11 (0.78 to 1.58)	1.46 (1.06 to 2.01)	1.45 (1.05 to 1.99)	0.94 (0.69 to 1.27) 1.1 (0.8 to 1.49)	1.58 (1.16 to 2.16)	1.61 (1.18 to 2.2)	1.13 (0.84 to 1.52)	1	
Natalizumab	0.3 (0.24 to 0.36)	1.03 (0.79 to 1.37)	1.88 (1.14 to 3.09)	0.43 (0.33 to 0.56)	0.36 (0.29 to 0.45)	0.47 (0.37 to 0.59)	0.35 (0.26 to 0.46)	0.46 (0.36 to 0.57)	0.45 (0.34 to 0.61)	0.59 (0.45 to 0.77)	0.59 (0.45 to 0.77)	0.39 (0.3 to 0.49) 0.45 (0.35 to 0.56	0.65 (0.5 to 0.83)	0.66 (0.51 to 0.84)	0.46 (0.33 to 0.25)	0.41 (0.29 to 0.57)	
Interferon beta-1b 250 mcg SC every other day	0.66 (0.57 to 0.76)	2.28 (1.84 to 2.94)	4.15 (2.6 to 6.71)	0.95 (0.77 to 1.19)	0.81 (0.69 to 0.93)	1.03 (0.88 to 1.22)	0.77 (0.61 to 0.96)	1.01 (0.88 to 1.16)	1.01 (0.78 to 1.3)	1.32 (1.06 to 1.65)	1.32 (1.06 to 1.63)	0.86 (0.69 to 1.03) 0.99 (0.81 to 1.2)	1.44 (1.17 to 1.75)	1.46 (1.19 to 1.78)	1.02 (0.75 to 0.83)	0.91 (0.66 to 1.22) 2.22 (1.76 to 2.81)	1
Interferon beta-1b 500 mcg SC every other day	0.62 (0.51 to 0.74)	2.13 (1.67 to 2.84)	3.87 (2.39 to 6.41)	0.89 (0.69 to 1.15)	0.76 (0.62 to 0.91)	0.96 (0.79 to 1.19)	0.72 (0.55 to 0.93)	0.95 (0.8 to 1.12)	0.94 (0.7 to 1.26)	1.24 (0.96 to 1.59)	1.23 (0.96 to 1.57)	0.8 (0.62 to 1.) 0.93 (0.73 to 1.17	1.34 (1.06 to 1.69)	1.37 (1.07 to 1.72)	0.96 (0.69 to 0.91)	0.85 (0.61 to 1.17) 2.07 (1.6 to 2.73)	0.93 (0.8 to 1.1)

# A6.2: Disability progression

Treatment	Placebo	Alemtuzumab 12 mg IV q.d	Alemtuzumab 24 mg IV q.d		a Interferon beta-1a 30 mcg IM q.w				240 mg two times	e dimethyl fumarate s 240 mg three times daily		al Teriflunomide oral 14 mg	al Fingolimod oral 0.5 mg	Fingolimod oral 1.25 mg	Peginterferon beta-1a 125 mcg once every 2 weeks	peginterferon beta-1a 125 mcg once every 4 weeks	Natalizumab	Interferon beta-1b 250 mcg SC every other day	
Placebo	1				,										,,	· · · · ·	· · · · · · · · · · · · · · · · · · ·		· · · · · · · · · · · · · · · · · · ·
Alemtuzumab 12 mg IV q.d	0.4 (0.27 to 0.6)	1		· [ · · · · · · · · · · · · · · · · · ·												· · · · · · · · · · · · · · · · · · ·			
Alemtuzumab 24 mg IV q.d		0.91 (0.42 to 1.8)	1	· [ · · · · · · · · · · · · · · · · · ·		,			· · · · · · · · · · · · · · · · · · ·						· · · · · · · · · · · · · · · · · · ·		1		
Interferon beta-1a 22 mcg SC t.i.w	0.84 (0.61 to 1.19)		-	J 1		,			1						,	· · · · · · · · · · · · · · · · · · ·	,,		1
Interferon beta-1a 30 mcg IM q.w			) 2.21 (1.05 to 4.94)		.j 1	· [ · · · · · · · · · · · · · · · · · ·			· · · · · · · · · · · · · · · · · · ·						· · · · · · · · · · · · · · · · · · ·	[	1		
Interferon beta-1a 44 mcg SC t.i.w			e) 2.15 (1.1 to 4.55)			1			,				1		,	· · · · · · · · · · · · · · · · · · ·			
Interferon beta-1a 60 mcg IM q.w					5) 0.99 (0.71 to 1.39)		.) 1		1				1		1	· · · · · · · · · · · · · · · · · · ·		1	
1.1	0.78 (0.63 to 0.96)	1			,	/		17) 1	· [ · · · · · · · · · · · · · · · · · ·						· [		1		
dimethyl fumarate 240 mg two	0.65 (0.49 to 0.85)								.) _ 1						· '		1'		
dimethyl fumarate 240 mg three times daily	0.68 (0.52 to 0.89)	1.73 (1.06 to 2.69)	) 1.9 (0.88 to 4.31)	0.81 (0.52 to 1.22	.) 0.86 (0.62 to 1.17	0.89 (0.61 to 1.26)	) 0.86 (0.54 to 1.3	6) 0.88 (0.64 to 1.18	.) 1.06 (0.78 to 1.42	.) 1					· · · · · · · · · · · · · · · · · · ·				
Teriflunomide oral 7 mg	0.8 (0.55 to 1.13)	2. (1.15 to 3.35)	2.2 (0.99 to 5.4)	0.94 (0.57 to 1.52)	2) 0.99 (0.65 to 1.48)	1.03 (0.65 to 1.58)	1.01 (0.58 to 1.6°	J) 1.01 (0.67 to 1.53	) 1.23 (0.78 to 1.91)	) 1.16 (0.74 to 1.81'	.1 1	<u> </u>	′	· [ ]	· · · · · · · · · · · · · · · · · · ·	<u> </u>			· [
Teriflunomide oral 14 mg	0.73 (0.51 to 1.05)	1.85 (1.06 to 3.11)	1 2.03 (0.91 to 4.89	J 0.87 (0.52 to 1.41'	L) 0.93 (0.6 to 1.38)	0.95 (0.6 to 1.46)	, 0.92 (0.54 to 1.5"	7) 0.94 (0.62 to 1.43	) 1.13 (0.72 to 1.76)	) 1.07 (0.68 to 1.69	1 0.92 (0.64 to 1.35'	.j 1			· · · · · · · · · · · · · · · · · · ·	· · · · ·			1
Fingolimod oral 0.5 mg	0.71 (0.55 to 0.9)	1.78 (1.11 to 2.77'	1.96 (0.92 to 4.55)	) 0.83 (0.54 to 1.26	5) 0.89 (0.65 to 1.16)	) 0.91 (0.64 to 1.3)	, 0.89 (0.57 to 1.3"	7) 0.9 (0.66 to 1.23'	1.09 (0.75 to 1.57	) 1.03 (0.72 to 1.48	) 0.89 (0.58 to 1.37'	) 0.97 (0.62 to 1.5)	1 1		· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·	1		
Fingolimod oral 1.25 mg	0.71 (0.56 to 0.9)	1.8 (1.12 to 2.78)	1.96 (0.93 to 4.49	) 0.85 (0.55 to 1.26	5) 0.89 (0.66 to 1.18)	) 0.92 (0.65 to 1.3)	0.9 (0.57 to 1.38	s) 0.91 (0.67 to 1.25	) 1.1 (0.77 to 1.59)	1.04 (0.73 to 1.49	) 0.9 (0.59 to 1.38)	, 0.97 (0.63 to 1.5)	1.01 (0.78 to 1.32	.j 1	· · · · · · · · · · · · · · · · · · ·				
Peginterferon beta-1a 125 mcg once every 2 weeks	0.61 (0.36 to 0.98)	1.53 (0.8 to 2.87)	1.68 (0.69 to 4.18	) 0.72 (0.39 to 1.28	) 0.76 (0.43 to 1.27	) 0.78 (0.44 to 1.36	) 0.77 (0.4 to 1.4'	) 0.78 (0.45 to 1.37	.) 0.94 (0.53 to 1.62	.) 0.89 (0.5 to 1.55'	0.77 (0.41 to 1.4)	0.83 (0.44 to 1.49	9) 0.86 (0.49 to 1.49)	) 0.86 (0.49 to 1.47	) 1		· '		
peginterferon beta-1a 125 mcg once every 4 weeks	0.62 (0.38 to 1.01)	1.56 (0.84 to 2.86	) 1.74 (0.71 to 4.23	) 0.73 (0.4 to 1.33)	0.78 (0.46 to 1.33	) 0.8 (0.46 to 1.38)	0.79 (0.41 to 1.4	4) 0.8 (0.47 to 1.37	, 0.96 (0.55 to 1.66	) 0.91 (0.52 to 1.57	) 0.79 (0.43 to 1.43	) 0.85 (0.46 to 1.55	5) 0.88 (0.51 to 1.54)	) 0.87 (0.51 to 1.53	) 1.02 (0.61 to 1.74	1			
Natalizumab	0.59 (0.42 to 0.84)	1.49 (0.86 to 2.5)	) 1.65 (0.73 to 3.9)	0.7 (0.43 to 1.13)	, 0.74 (0.49 to 1.11'	) 0.76 (0.49 to 1.18'	) 0.75 (0.44 to 1.2"	5) 0.75 (0.5 to 1.15'	, 0.91 (0.59 to 1.42'	.) 0.86 (0.56 to 1.34	) 0.74 (0.45 to 1.23	.) 0.8 (0.49 to 1.34)	) 0.84 (0.55 to 1.28)	) 0.83 (0.55 to 1.27	) 0.97 (0.53 to 1.81'	0.94 (0.53 to 1.73)	/ 1		
	0.72 (0.54 to 0.92)	1.8 (1.1 to 2.77)	1.97 (0.92 to 4.52	) 0.85 (0.54 to 1.26	) 0.9 (0.65 to 1.17)	0.93 (0.64 to 1.28	) 0.9 (0.56 to 1.36	) 0.92 (0.69 to 1.1f	) 1.1 (0.75 to 1.58)	, 1.05 (0.72 to 1.48	.) 0.9 (0.57 to 1.38)	, 0.98 (0.61 to 1.51	1) 1.02 (0.7 to 1.42)	, 1.01 (0.69 to 1.41	.) 1.17 (0.66 to 2.07	1.16 (0.63 to 0.74)	1.22 (0.76 to 1.84	,) 1	
Interferon beta-1b 500 mcg SC every other day	0.79 (0.56 to 1.1)	1.99 (1.18 to 3.2)	2.18 (1. to 5.02)	0.94 (0.58 to 1.47	) 0.99 (0.68 to 1.39	1.02 (0.68 to 1.49)	) 1. (0.6 to 1.59)	1.01 (0.74 to 1.3f	) 1.22 (0.8 to 1.84)	, 1.15 (0.77 to 1.73	.) 1. (0.6 to 1.61)	1.08 (0.65 to 1.76	6) 1.12 (0.74 to 1.68)	) 1.11 (0.73 to 1.66	) 1.3 (0.71 to 2.38)	1.27 (0.68 to 1.19)	, 1.34 (0.81 to 2.16)	) 1.1 (0.84 to 1.51)	, <u>1</u>

# A6.3: Withdrawal due to adverse events

Treatment	Placebo	Alemtuzumab 12 mg IV q.d	Alemtuzumab 24 mg IV q.d		Interferon beta-1a 30 mcg IM q.w			glatiramer acetate 20mg q.d	glatiramer acetate 40mg t.i.w	240 mg two times	dimethyl fumarate 240 mg three times daily		l Teriflunomide oral 14 mg	Fingolimod oral 0.5 mg	Fingolimod oral 1.25 mg	Peginterferon beta-1a 125 mcg once every 2 weeks	peginterferon beta-1a 125 mcg once every 4 weeks	Natalizumab	Interferon beta 1b 250 mcg SC every other day	every other
Placebo	1			-		-						-	-	_	_					
Alemtuzumab 12 mg IV q.d	0.61 (0.25 to 1.47)	1																		
Alemtuzumab 24 mg IV q.d	0.54 (0.17 to 1.54)	0.88 (0.3 to 2.31)	1																	
Interferon beta-1a 22 mcg SC t.i.w	1.68 (0.5 to 5.98)	2.78 (0.7 to 11.12)	3.16 (0.7 to 15.17)	1																
Interferon beta-1a 30 mcg IM q.w	1.33 (0.85 to 2.17)	2.18 (0.89 to 5.5)	2.46 (0.85 to 8.1)	0.8 (0.22 to 2.82)	1															
Interferon beta-1a 44 mcg SC t.i.w	2.2 (1.29 to 3.97)	3.6 (1.88 to 7.33)	4.08 (1.69 to 11.42	) 1.31 (0.4 to 4.36)	1.65 (0.91 to 3.08)	1														
Interferon beta-1a 60 mcg IM q.w	1.9 (0.79 to 4.81)	3.1 (0.96 to 10.5)	3.5 (0.95 to 14.59)	1.14 (0.25 to 4.94)	1.43 (0.66 to 3.11)	0.86 (0.32 to 2.29	1													
glatiramer acetate 20mg q.d	1.17 (0.74 to 1.94)	1.91 (0.79 to 4.89)	2.16 (0.76 to 7.2)	0.7 (0.19 to 2.48)	0.88 (0.51 to 1.55)	0.53 (0.29 to 0.96	0.62 (0.24 to 1.63	1												
glatiramer acetate 40mg t.i.w	2.5 (0.86 to 8.29)	4.08 (1.02 to 18.4)	4.7 (1.05 to 24.1)	1.47 (0.29 to 8.02)	1.87 (0.58 to 6.69)	1.13 (0.33 to 4.18	1.32 (0.32 to 5.79	2.13 (0.65 to 7.5)	1											
dimethyl fumarate 240 mg two times daily	1.24 (0.74 to 2.13)	2.02 (0.75 to 5.59)	2.29 (0.74 to 8.12)	0.74 (0.19 to 2.73)	0.94 (0.47 to 1.82)	0.56 (0.27 to 1.15	0.66 (0.23 to 1.81	1.07 (0.56 to 1.92)	0.5 (0.14 to 1.64)	1										
dimethyl fumarate 240 mg three times daily	1.25 (0.74 to 2.13)	2.03 (0.76 to 5.6)	2.32 (0.74 to 8.16)	0.75 (0.19 to 2.74)	0.94 (0.47 to 1.83)	0.57 (0.27 to 1.17	0.66 (0.23 to 1.81	) 1.07 (0.56 to 1.93)	0.5 (0.14 to 1.65)	1.01 (0.58 to 1.73)	1									
Teriflunomide oral 7 mg	1.37 (0.82 to 2.21)	2.24 (0.87 to 5.55)	2.53 (0.85 to 8.25)	0.82 (0.22 to 2.84)	1.03 (0.52 to 1.91)	0.62 (0.31 to 1.12	0.72 (0.25 to 1.9)	1.17 (0.57 to 2.16)	0.55 (0.15 to 1.74)	1.1 (0.52 to 2.19)	1.1 (0.52 to 2.19)	1								
Teriflunomide oral 14 mg	1.53 (0.96 to 2.54)	2.51 (1.02 to 6.37)	2.85 (0.98 to 9.45)	0.9 (0.25 to 3.25)	1.15 (0.61 to 2.19)	0.69 (0.37 to 1.28	0.81 (0.29 to 2.2)	1.31 (0.68 to 2.48)	0.62 (0.17 to 1.99)	1.23 (0.62 to 2.54)	1.23 (0.61 to 2.53)	1.12 (0.73 to 1.85	) 1							
Fingolimod oral 0.5 mg	1.54 (0.98 to 2.52)	2.52 (0.96 to 6.8)	2.85 (0.93 to 9.82)	0.91 (0.24 to 3.35)	1.16 (0.65 to 2.04)	0.7 (0.34 to 1.4)	0.81 (0.31 to 2.1)	1.31 (0.68 to 2.48)	0.62 (0.17 to 1.99)	1.24 (0.62 to 2.52)	1.24 (0.62 to 2.5)	1.12 (0.59 to 2.29	1.01 (0.52 to 1.96)	1						
Fingolimod oral 1.25 mg	2.21 (1.42 to 3.58)	3.62 (1.38 to 9.71)	4.09 (1.34 to 14.02	) 1.31 (0.35 to 4.8)	1.66 (0.94 to 2.91)	1. (0.49 to 1.99)	1.16 (0.45 to 3.02	1.88 (0.99 to 3.53)	0.89 (0.25 to 2.84)	1.78 (0.89 to 3.61)	1.77 (0.89 to 3.6)	1.6 (0.86 to 3.27)	1.45 (0.74 to 2.79)	1.43 (0.94 to 2.21)	1					
Peginterferon beta-1a 125 mcg once every 2 weeks	3.57 (1.27 to 11.14)	5.78 (1.51 to 24.42	) 6.67 (1.5 to 33.69)	2.12 (0.41 to 11.18	) 2.67 (0.85 to 9.09)	1.62 (0.49 to 5.69	1.88 (0.47 to 7.82	3.04 (0.95 to 10.19	1.43 (0.29 to 6.88)	2.87 (0.9 to 9.95)	2.85 (0.9 to 9.93)	2.59 (0.84 to 9.15	2.31 (0.73 to 8.02)	2.3 (0.73 to 7.85)	1.61 (0.51 to 5.43)	1				
peginterferon beta-1a 125 mcg once every 4 weeks	3.47 (1.25 to 10.9)	5.75 (1.48 to 24.35	) 6.48 (1.48 to 33.07	) 2.07 (0.4 to 10.92)	2.61 (0.83 to 8.91)	1.58 (0.48 to 5.61	1.83 (0.46 to 7.75	2.96 (0.94 to 10.)	1.4 (0.29 to 6.78)	2.8 (0.87 to 9.8)	2.78 (0.88 to 9.76)	2.54 (0.83 to 8.99	2.28 (0.72 to 7.88)	2.27 (0.72 to 7.71)	1.59 (0.5 to 5.33)	0.98 (0.41 to 2.37)	1			
Natalizumab	1.22 (0.5 to 2.74)	1.98 (0.56 to 6.49)	2.26 (0.56 to 8.94)	0.72 (0.15 to 3.06)	0.91 (0.32 to 2.31)	0.55 (0.18 to 1.44	0.65 (0.17 to 2.1)	1.04 (0.36 to 2.6)	0.48 (0.11 to 1.84)	0.98 (0.34 to 2.52)	0.98 (0.33 to 2.53)	0.89 (0.32 to 2.3)	0.79 (0.28 to 2.)	0.79 (0.28 to 1.98)	0.55 (0.19 to 1.36)	0.34 (0.08 to 1.54)	0.34 (0.08 to 1.28)	1		
Interferon beta-1b 250 mcg SC every other day	0.84 (0.4 to 1.87)	1.36 (0.46 to 4.29)	1.56 (0.46 to 6.11)	0.49 (0.12 to 2.07)	0.63 (0.28 to 1.44)	0.38 (0.16 to 0.93	0.44 (0.15 to 1.37	0.72 (0.35 to 1.49)	0.33 (0.09 to 1.28)	0.68 (0.29 to 1.67)	0.67 (0.29 to 1.67)	0.61 (0.26 to 1.57	0.55 (0.23 to 1.34)	0.54 (0.23 to 1.32)	0.38 (0.16 to 0.92)	0.23 (0.06 to 5.98)	0.24 (0.06 to 0.9)	0.7 (0.23 to 2.37)	1	
Interferon beta-1b 500 mcg SC every other day	1.37 (0.52 to 3.92)	2.25 (0.63 to 8.47)	2.55 (0.64 to 11.48	) 0.8 (0.17 to 3.91)	1.03 (0.37 to 2.99)	0.62 (0.21 to 1.9)	0.72 (0.2 to 2.71)	1.16 (0.46 to 3.05)	0.54 (0.12 to 2.46)	1.1 (0.38 to 3.38)	1.09 (0.38 to 3.39)	1. (0.34 to 3.22)	0.89 (0.3 to 2.74)	0.89 (0.3 to 2.73)	0.62 (0.21 to 1.9)	0.38 (0.09 to 2.17)	0.39 (0.09 to 1.75)	1.13 (0.32 to 4.63	63 (0.66 to 4.1	1 1

# A6.4: Change in Expanded Disability Status Scale

Treatment	Placebo	Alemtuzumab 12 mg IV q.d	Alemtuzumab 24 mg IV q.d	Interferon beta-1a 22 mcg SC t.i.w	Interferon beta-1a 30 mcg IM q.w			glatiramer acetate 20mg q.d		Teriflunomide oral 14 mg	Fingolimod oral 0.5 mg	Fingolimod oral 1.25 mg	Interferon beta-1b 250 mcg SC every other day
Placebo	1												
Alemtuzumab 12 mg IV q.d	-0.6 (-1.02 to -0.24)	1											
Alemtuzumab 24 mg IV q.d	-0.91 (-1.48 to -0.4)	-0.31 (-0.76 to 0.15)	1										
Interferon beta-1a 22 mcg SC t.i.w	-0.27 (-0.71 to 0.15)	0.33 (-0.15 to 0.85)	0.64 (0.03 to 1.28)	1									
Interferon beta-1a 30 mcg IM q.w	-0.22 (-0.48 to 0.02)	0.38 (0.04 to 0.77)	0.69 (0.18 to 1.24)	0.05 (-0.4 to 0.51)	1								
Interferon beta-1a 44 mcg SC t.i.w	-0.28 (-0.58 to -0.02)		0.63 (0.18 to 1.1)	-0.01 (-0.44 to 0.41)	-0.06 (-0.32 to 0.18	) 1							
Interferon beta-1a 60 mcg IM q.w	-0.25 (-0.76 to 0.24)		0.66 (0. to 1.36)			0.03 (-0.46 to 0.54)	1						
glatiramer acetate 20mg q.d	-0.13 (-0.4 to 0.11)	0.47 (0.08 to 0.9)	0.78 (0.24 to 1.35)			0.15 (-0.15 to 0.47)		1					
Teriflunomide oral 7 mg	-0.05 (-0.47 to 0.36)	0.55 (0.01 to 1.15)	0.86 (0.21 to 1.57)	0.22 (-0.36 to 0.83)	0.17 (-0.3 to 0.66)	0.23 (-0.25 to 0.75)	0.19 (-0.44 to 0.86	0.08 (-0.4 to 0.58)	1				
Teriflunomide oral 14 mg	-0.14 (-0.56 to 0.27)			0.13 (-0.46 to 0.74)						1			
Fingolimod oral 0.5 mg	-0.16 (-0.41 to 0.1)	0.44 (0.04 to 0.91)	0.76 (0.21 to 1.36)	0.12 (-0.36 to 0.61)	0.06 (-0.22 to 0.36)	0.12 (-0.2 to 0.48)	0.09 (-0.42 to 0.63	-0.03 (-0.35 to 0.33	) -0.1 (-0.59 to 0.38)	-0.02 (-0.5 to 0.47)	1		
Fingolimod oral 1.25 mg	-0.22 (-0.47 to 0.04)										-0.06 (-0.3 to 0.18)	1	
Interferon beta-1b 250 mcg SC every other day	-0.58 (-0.94 to -0.22)												) 1

## A6.5: Serious adverse events

Treatment	Placebo	Alemtuzumab 12 mg IV q.d	Alemtuzumab 24 mg IV q.d		Interferon beta-1a 44 mcg SC t.i.w	0	0	dimethyl fumarate 240 mg two times daily	,		l Teriflunomide oral 14 mg	Fingolimod oral 0.5 mg	Fingolimod oral 1.25 mg	Peginterferon beta-1a 125 mcg once every 2 weeks	peginterferon beta-1a 125 mcg once every 4 weeks	Natalizumab		Interferon beta-1b 500 mcg SC every other day
Placebo	1																	
Alemtuzumab 12 mg IV q.d	0.67 (0.37 to 1.28)	1																
Alemtuzumab 24 mg IV q.d	0.79 (0.42 to 1.53)	1.18 (0.79 to 1.71)	1															
Interferon beta-1a 30 mcg IM q.w	0.77 (0.54 to 1.13)	1.14 (0.61 to 2.07)	0.97 (0.51 to 1.83)	1														
Interferon beta-1a 44 mcg SC t.i.w	0.86 (0.52 to 1.46)	1.28 (0.91 to 1.75)	1.09 (0.74 to 1.59)	1.12 (0.67 to 1.86)	1													
glatiramer acetate 20mg q.d	0.78 (0.54 to 1.14)	1.16 (0.62 to 2.08)	0.99 (0.52 to 1.83)	1.01 (0.67 to 1.53)	0.91 (0.55 to 1.49)	1												
glatiramer acetate 40mg t.i.w	0.99 (0.49 to 2.04)	1.47 (0.57 to 3.72)	1.25 (0.48 to 3.23)	1.28 (0.58 to 2.87)	1.15 (0.48 to 2.75)	1.27 (0.57 to 2.83)	1											
dimethyl fumarate 240 mg two times daily	0.81 (0.56 to 1.19)	1.21 (0.59 to 2.36)	1.03 (0.49 to 2.07)	1.05 (0.63 to 1.72)	0.94 (0.51 to 1.71)	1.04 (0.65 to 1.63)	0.82 (0.36 to 1.81	1										
dimethyl fumarate 240 mg three times daily	0.72 (0.49 to 1.07)	1.08 (0.52 to 2.1)	0.92 (0.44 to 1.84)	0.94 (0.56 to 1.54)	0.84 (0.45 to 1.53)	0.93 (0.58 to 1.46)	0.73 (0.33 to 1.62	0.89 (0.6 to 1.33)	1									
Teriflunomide oral 7 mg	1.03 (0.71 to 1.51)	1.54 (0.77 to 2.92)	1.31 (0.64 to 2.58)	1.34 (0.8 to 2.2)	1.2 (0.67 to 2.12)	1.33 (0.79 to 2.19)	1.05 (0.46 to 2.31	1.28 (0.75 to 2.16)	1.43 (0.84 to 2.44)	1								
Teriflunomide oral 14 mg	1.07 (0.73 to 1.54)	1.58 (0.78 to 3.01)	1.35 (0.66 to 2.65)	1.38 (0.82 to 2.26)	1.24 (0.68 to 2.18)	1.37 (0.81 to 2.24)	1.08 (0.48 to 2.36	1.32 (0.77 to 2.21)	1.48 (0.86 to 2.49)	1.03 (0.71 to 1.48)	) 1							
Fingolimod oral 0.5 mg	0.96 (0.68 to 1.39)	1.43 (0.71 to 2.77)	1.22 (0.59 to 2.44)	1.25 (0.8 to 1.95)	1.12 (0.61 to 2.01)	1.24 (0.76 to 2.01)	0.97 (0.44 to 2.14	1.19 (0.71 to 1.99)	1.33 (0.8 to 2.25)	0.93 (0.56 to 1.56)	0.9 (0.55 to 1.52)	1						
Fingolimod oral 1.25 mg	1.22 (0.87 to 1.77)	1.81 (0.91 to 3.53)	1.54 (0.76 to 3.11)	1.58 (1.03 to 2.47)	1.41 (0.79 to 2.56)	1.56 (0.97 to 2.55)	1.23 (0.56 to 2.74	1.5 (0.92 to 2.55)	1.68 (1.03 to 2.88)	1.18 (0.72 to 1.99)	1.14 (0.7 to 1.95)	1.26 (0.91 to 1.8)	1					
Peginterferon beta-1a 125 mcg once every 2 weeks	1.67 (0.94 to 2.94)	2.48 (1.04 to 5.55)	2.11 (0.88 to 4.9)	2.16 (1.08 to 4.21)	1.93 (0.88 to 4.1)	2.14 (1.06 to 4.16)	1.69 (0.68 to 4.13	2.06 (1.03 to 4.07)	2.31 (1.16 to 4.57)	1.62 (0.81 to 3.16)	) 1.56 (0.79 to 3.1)	1.73 (0.87 to 3.34)	1.37 (0.69 to 2.62)	1				
peginterferon beta-1a 125 mcg once every 4 weeks	1.55 (0.88 to 2.74)	2.31 (0.97 to 5.19)	1.96 (0.81 to 4.57)	2.02 (1. to 3.95)	1.8 (0.82 to 3.85)	2. (0.99 to 3.89)	1.57 (0.63 to 3.84	1.92 (0.95 to 3.8)	2.15 (1.07 to 4.25)	1.5 (0.75 to 2.96)	1.45 (0.74 to 2.9)	1.61 (0.81 to 3.12)	1.28 (0.64 to 2.43)	0.93 (0.54 to 1.61)	1			
Natalizumab	0.81 (0.49 to 1.39)	1.21 (0.53 to 2.62)	1.03 (0.45 to 2.31)	1.06 (0.56 to 1.99)	0.95 (0.46 to 1.93)	1.04 (0.56 to 1.95)	0.82 (0.34 to 1.97	1.01 (0.53 to 1.92)	1.13 (0.6 to 2.15)	0.79 (0.42 to 1.5)	0.76 (0.41 to 1.47)	0.85 (0.46 to 1.59)	0.67 (0.36 to 1.23)	0.49 (0.23 to 1.07)	0.52 (0.25 to 1.15)	1		
Interferon beta-1b 250 mcg SC every other day	0.66 (0.35 to 1.26)	0.99 (0.43 to 2.18)	0.84 (0.36 to 1.9)	0.86 (0.43 to 1.68)	0.77 (0.36 to 1.61)	0.85 (0.49 to 1.45)	0.67 (0.25 to 1.72)	0.82 (0.4 to 1.64)	0.92 (0.45 to 1.86)	0.64 (0.31 to 1.33)	) 0.62 (0.3 to 1.31)	0.69 (0.33 to 1.41)	0.55 (0.26 to 1.09)	0.4 (0.17 to 0.94)	0.43 (0.18 to 1.02)	0.82 (0.36 to 1.78)	1	
Interferon beta-1b 500 mcg SC every other day	0.93 (0.49 to 1.8)	1.38 (0.6 to 3.05)	1.18 (0.5 to 2.67)	1.21 (0.61 to 2.36)	1.08 (0.51 to 2.25)	1.19 (0.69 to 2.06)	0.94 (0.36 to 2.43	1.15 (0.57 to 2.33)	1.29 (0.64 to 2.63)	0.9 (0.43 to 1.9)	0.87 (0.42 to 1.85)	0.97 (0.46 to 2.)	0.77 (0.36 to 1.55)	0.56 (0.24 to 1.34)	0.6 (0.25 to 1.44)	1.14 (0.5 to 1.53)	1.4 (0.83 to 2.4)	1

## A6.6: Mortality

Treatment	Placebo	Alemtuzumab 12 mg IV q.d	Alemtuzumab 24 mg IV q.d		Interferon beta-1a 30 mcg IM q.w			glatiramer acetate 20mg q.d	glatiramer acetate 40mg t.i.w	dimethyl fumarate 240 mg two times daily		Teriflunomide oral 7 mg	Teriflunomide oral 14 mg	Fingolimod oral 0.5 mg	Fingolimod oral 1.25 mg	Peginterferon beta-1a 125 mcg once every 2 weeks	peginterferon beta-1a 125 mcg once every 4 weeks	Natalizumab	Interferon beta- 1b 250 mcg SC every other day	every other
Placebo	1																			
Alemtuzumab 12 mg IV q.d	2.81 (0.08 to 168.2)	1																		
Alemtuzumab 24 mg IV q.d	2.08 (0.04 to 125.5)	0.73 (0.06 to 5.88)	1																	
Interferon beta-1a 22 mcg SC t.i.w	1.6 (0.07 to 34.77)	0.55 (0.01 to 24.67)	0.78 (0.01 to 51.33	1																
Interferon beta-1a 30 mcg IM q.w	2.1 (0.26 to 24.45)	0.82 (0.01 to 40.09	1.09 (0.01 to 84.65	1.4 (0.04 to 55.3)	1															
Interferon beta-1a 44 mcg SC t.i.w	0.97 (0.06 to 17.15)				0.43 (0.01 to 12.35)	1														
Interferon beta-1a 60 mcg IM q.w	2.28 (0.03 to 222.1)	0.88 (0. to 177.5)	1.15 (0. to 358.5)	1.51 (0.01 to 308.9	1.01 (0.02 to 55.64)	2.48 (0.01 to 420.6)	1													
glatiramer acetate 20mg q.d	0.9 (0.11 to 7.85)	0.33 (0.01 to 10.23)	0.44 (0.01 to 19.73	0.55 (0.02 to 22.44	0.42 (0.03 to 4.39)	0.97 (0.06 to 14.14)	0.41 (0. to 42.93)	1												
glatiramer acetate 40mg t.i.w	0.08 (0. to 3.54)				0.04 (0. to 2.97)			0.09 (0. to 7.16)	1											
dimethyl fumarate 240 mg two times daily	0.52 (0.04 to 5.34)	0.18 (0. to 10.34)	0.24 (0. to 19.9)	0.32 (0.01 to 14.89	0.24 (0.01 to 4.56)	0.51 (0.02 to 16.38)	0.22 (0. to 29.21)	0.57 (0.03 to 7.89)	6.69 (0.06 to 7441.	1										
dimethyl fumarate 240 mg three times daily	0.89 (0.09 to 8.41)	0.3 (0. to 16.87)	0.42 (0. to 32.89)	0.53 (0.01 to 25.59	) 0.42 (0.02 to 7.26)	0.9 (0.03 to 26.01)	0.4 (0. to 46.87)	0.98 (0.08 to 11.73	1.07 (0.13 to 11810	1.69 (0.18 to 18.19)	1									
Teriflunomide oral 7 mg	2.59 (0.12 to 82.51)	0.93 (0. to 126.9)	1.36 (0. to 242.8)	1.66 (0.02 to 167.7	) 1.16 (0.03 to 67.28)	2.73 (0.04 to 238.5)	.08 (0.01 to 383.4	2.88 (0.07 to 196.5	6.05 (0.27 to 53180	5.18 (0.12 to 307.1)	3.08 (0.07 to 174.6	1								
Teriflunomide oral 14 mg	0.94 (0.02 to 37.74)	0.33 (0. to 56.46)	0.48 (0. to 98.06)	0.58 (0. to 88.85)	0.45 (0. to 34.11)	0.99 (0.01 to 105.1)	0.42 (0. to 157.2)	1.05 (0.01 to 71.67	2.64 (0.05 to 19540	1.88 (0.02 to 147.7)	1.1 (0.01 to 79.34)	0.39 (0.01 to 7.5)	1							
Fingolimod oral 0.5 mg	0.1 (0. to 2.57)	0.03 (0. to 4.24)	0.04 (0. to 8.67)	0.05 (0. to 5.39)	0.04 (0. to 2.2)	0.08 (0. to 8.4)	0.04 (0. to 12.77)	0.1 (0. to 5.12)	1.03 (0. to 1842.)	0.17 (0. to 12.01)	0.1 (0. to 6.16)	0.03 (0. to 3.64)	0.09 (0. to 18.86)	1						
Fingolimod oral 1.25 mg	0.52 (0.02 to 6.76)	0.17 (0. to 13.59)	0.24 (0. to 24.79)	0.31 (0. to 17.09)	0.24 (0.01 to 6.36)	0.5 (0.01 to 23.87)	0.22 (0. to 40.74)	0.57 (0.01 to 15.09	6.57 (0.05 to 8530.	0.98 (0.02 to 36.2)	0.59 (0.01 to 17.31	0.19 (0. to 9.23)	0.5 (0. to 63.23)	5.46 (0.12 to 4103.)	1					
Peginterferon beta-1a 125 mcg once every 2 weeks	0.41 (0.01 to 8.87)	0.13 (0. to 12.73)	0.18 (0. to 25.17)	0.25 (0. to 19.58)	0.17 (0. to 8.06)	0.41 (0. to 22.98)	0.17 (0. to 38.09)	0.43 (0.01 to 19.38	5.11 (0.02 to 7408.	0.79 (0.01 to 44.58)	0.44 (0.01 to 21.56	0.15 (0. to 11.05)	0.42 (0. to 71.51)	4.72 (0.02 to 4890.)	0.79 (0.01 to 65.78)	1				
peginterferon beta-1a 125 mcg once every 4 weeks	0.4 (0.01 to 10.22)	0.13 (0. to 14.86)	0.18 (0. to 26.14)	0.24 (0. to 19.82)	0.17 (0. to 8.86)	0.39 (0. to 24.78)	0.16 (0. to 42.68)	0.42 (0. to 20.5)	4.75 (0.02 to 7254.	0.78 (0.01 to 43.94)	0.45 (0. to 21.3)	0.14 (0. to 10.48)	0.4 (0. to 69.74)	4.66 (0.03 to 5197.	0.74 (0.01 to 66.19)	1. (0.02 to 44.74)	1			
Natalizumab	4.34 (0.16 to 2761.)	1.73 (0.01 to 2475.)	2.19 (0.01 to 4596.)	3.03 (0.04 to 3566.	) 2.21 (0.03 to 2092.)	5.17 (0.06 to 4779.)	.53 (0.01 to 3792.	5.25 (0.1 to 4005.)	3.25 (0.34 to 20170	9.32 (0.14 to 8262.)	5.45 (0.09 to 4151.)	1.88 (0.02 to 1428.)	5.75 (0.03 to 5626.)	5.43 (0.48 to 34030	9.91 (0.14 to 10330.	12.92 (0.12 to 125.5	3.91 (0.12 to 23380	1		
Interferon beta-1b 250 mcg SC every other day	0.07 (0. to 6.65)				0.03 (0. to 3.51)														1	
Interferon beta-1b 500 mcg SC every other day	0.08 (0. to 5.9)	0.02 (0. to 4.32)	0.04 (0. to 8.01)	0.05 (0. to 8.57)	0.03 (0. to 3.12)	0.08 (0. to 7.76)	0.03 (0. to 13.35)	0.09 (0. to 3.55)	0.95 (0. to 2803.)	0.15 (0. to 17.67)	0.09 (0. to 8.48)	0.03 (0. to 5.52)	0.08 (0. to 31.17)	0.87 (0. to 1764.)	0.15 (0. to 31.31)	0.19 (0. to 24.45)	0.19 (0. to 128.)	0.01 (0. to 4.47)	1.08 (0. to 863.8	) 1

#### Appendix 7: Results for direct pairwise meta-analyses

Interventions	RR (95% CI)			
Natalizumab	0.31 (0.26 to 0.36)			
Fingolimod oral 1.25 mg	0.44 (0.38 to 0.51)			
Fingolimod oral 0.5 mg	0.45 (0.41 to 0.56)			
Dimethyl fumarate 240 mg two times daily	0.51 (0.44 to 0.60)			
Dimethyl fumarate 240 mg three times daily	0.51 (0.44 to 0.60)			
Interferon beta-1a 44 mcg SC t.i.w.	0.67 (0.58 to 0.77)			
Peg-interferon beta-1a 125 mcg once every 2 weeks	0.64 (0.50 to 0.82)			
Glatiramer acetate 20mg q.d.	0.71 (0.62 to 0.80)			
Glatiramer acetate 40mg t.i.w.	0.66 (0.55 to 0.78)			
Interferon beta-1b 250 mcg SC every other day	0.65 (0.54 to 0.79)			
Teriflunomide oral 14 mg	0.66 (0.58 to 0.75)			
Interferon beta-1a 22 mcg SC t.i.w.	0.71 (0.62 to 0.82)			
Peg-interferon beta-1a 125 mcg once every 4 weeks	0.73 (0.57 to 0.92)			
Teriflunomide oral 7 mg	0.73 (0.65 to 0.82)			
Interferon beta-1a 30 mcg IM q.w.	0.79 (0.69 to 0.89)			

#### A7.1 Annual relapse for multiple sclerosis treatments compared to placebo

RR= relative ratio, CI= confidence interval, mg= milligrams, mcg= micrograms, SC= subcutaneous, IM= intra muscular, q.d.= once daily, q.w.=once weekly, t.i.w.= three times weekly,

### A7.2: Disability progression for multiple sclerosis treatments compared to placebo

Interventions	RR (95% CI)
Natalizumab	0.59 (0.46, 0.75)
Peg-interferon beta-1a 125 mcg once every 2	
weeks	0.61 (0.39, 0.93)
Peg-interferon beta-1a 125 mcg once every 4	
weeks	0.62(0.40,0.95)
Dimethyl fumarate 240 mg two times daily	0.66 (0.52, 0.84)
Dimethyl fumarate 240 mg three times daily	0.70 (0.57, 0.86)
Fingolimod oral 0.5 mg	0.75(0.60,0.92)
Fingolimod oral 1.25 mg	0.70 (0.57, 0.87)
Interferon beta-1b 250 mcg SC every other day	0.77 (0.56, 1.04)
Teriflunomide oral 14 mg	0.74 (0.57, 0.96)
Interferon beta-1a 44 mcg SC t.i.w	0.71(0.54,0.95)
Glatiramer acetate 20mg q.d	0.92(0.70, 1.20)
Teriflunomide oral 7 mg	0.79 (0.61, 1.03)
Interferon beta-1a 30 mcg IM q.w	0.69 (0.52, 0.91)
Interferon beta-1a 22 mcg SC t.i.w	0.82 (0.63, 1.07)

RR= relative ratio, CI= confidence interval, mg= milligrams, mcg= micrograms, SC= subcutaneous, IM= intra muscular, q.d.= once daily, q.w.=once weekly, t.i.w.= three times weekly,

### A7.3: Withdrawal due to adverse events for multiple sclerosis treatments compared to placebo

Interventions	RR (95% CI)
Interferon beta-1b 250 mcg SC every other day	0.10 (0.01, 0.76)
Glatiramer acetate 20mg q.d	1.20 (0.59, 2.43)
Dimethyl fumarate 240 mg two times daily	1.21(0.93, 1.57)
Dimethyl fumarate 240 mg three times daily	1.18 (0.91, 1.53)
Interferon beta-1a 30 mcg IM q.w	1.55 (0.91, 2.65)
Teriflunomide oral 7 mg	1.54 (0.81, 2.94)
Teriflunomide oral 14 mg	1.70 (1.25, 2.33)
Fingolimod oral 0.5 mg	1.41 (0.89, 2.24)
Interferon beta-1a 22 mcg SC t.i.w	2.97 (0.61, 14.52)
Glatiramer acetate 40mg t.i.w	2.36 (0.99, 5.65)
Interferon beta-1a 44 mcg SC t.i.w	4.57 (1.00, 20.88)
Fingolimod oral 1.25 mg	1.87 (1.43, 2.45)
Peg-interferon beta-1a 125 mcg once every 4 weeks	3.43 (1.49, 7.88)
Peg-interferon beta-1a 125 mcg once every 2 weeks	3.49 (1.52, 7.99)

RR= relative ratio, CI= confidence interval, mg= milligrams, mcg= micrograms, SC= subcutaneous,

IM= intra muscular, q.d.= once daily, q.w.=once weekly, t.i.w.= three times weekly,

### Appendix 8 Monitorings costs

#### 8.1: Monitoring costs associated with each of the treatments (1. year)

Drug	NAB-analyses	Infusion costs	Eye examina- tions	Startup costs	Medical consulta- tions	MRI	Blood tests (outpatient visits)	Travel costs	Total
Alemtuzumab (Lemtrada)	0	9777 (5/year)	0	0	7350 (4/year)	1 (!/year)	1008 (9/year)	2000 a	22,735
Dimethyl fumarate (Tecifidera)	0	0	0	0	7350 (4/year)	2600 (1/year)	0	1600 ^b	11,550
Fingolimod (Gilenya)	0	0	2500 (1/year)	3750 °	7350 (4/year)	2600 (1/year)	112 (1/year)	1600 ^b	17,912
Glatiramer ace- tate (Copaxone)	0	0	0	0	7350 (4/year)	2600 (1/year)	0	1600 ^b	11,550
Interferon beta- 1a (Avonex)	7716 (2/year)	0	0	0	7350 (4/year)	2600 (1/year)	0	1600 ^b	19,266

Interferon beta- 1a 44 mcg (Rebif)	7716 (2/year)	0	0	0	7350 (4/year)	2600 (1/year)	0	1600 ^b	19,266
Interferon beta- 1a 22 mcg (Rebif)	7716 (2/year)	0	0	0	7350 (4/year)	2600 (1/year)	0	1600 ^b	19,266
Interferon beta- 1b (Betaferon)	7716 (2/year)	0	0	0	7350 (4/year)	2600 (1/year)	0	1600 ^b	19,266
Interferon beta- 1b (Extavia)	7716 (2/year)	0	0	0	7350 (4/year)	2600 (1/year)	0	1600 ^b	19,266
Natalizumab (Tysabri)	1840 (2/year)	16,250 (13/year)	0	0	7350 (4/year)	2600 (1/year)	0	5200 ª	33,240
Peg-interferon beta-1a (Plegridy)	7716 (2/year)	0	0	0	7350 (4/year)	2600 (1/year)	0	1600 ^b	19,266
Teriflunomide (Aubagio)	0	0	0	0	7350 (4/year)	2600 (1/year)	1344 d	1600 ^ь	12,894

^a Analyses, MR, medical consultations and infusions will be done at the same day.

^b Analyses, MR, and medical consultations will be done at the same day (4/year).

^c6 hours observation

^d Every 14 days for 6 months, then every other month (numbers of medical consultations were deducted)

8.2: Monitoring costs associated with each of the treatment	s (2. year)
-------------------------------------------------------------	-------------

Drug	NAB-analyses	Infusion costs	Eye examina- tions	Startup costs	Medical consulta- tions	MRI	Blood tests (outpatient visits)	Travel costs	Total
Alemtuzumab (Lemtrada)	0	5866 (3/year)	0	0	3675 (2/year)	2600 (1/year)	1232 (11/year)	1200 ª	14,573
Dimethyl fumarate (Tecifidera)	0	0	0	0	3675 (2/year)	2600 (1/year)	0	800 b	7075
Fingolimod (Gilenya)	0	0	0	0	3675 (2/year)	2600 (1/year)	0	800 b	7075
Glatiramer acetate (Copaxone)	0	0	0	0	3675 (2/year)	2600 (1/year)	0	800 b	7075
Interferon beta- 1a (Avonex)	7716 (2/year)	0	0	0	3675 (2/year)	2600 (1/year)	0	800 ^b	14,791
Interferon beta- 1a 44 mcg (Rebif)	7716 (2/year)	0	0	0	3675 (2/year)	2600 (1/year)	0	800 b	14,791

Interferon beta- 1a 22 mcg (Rebif)	7716 (2/year)	0	0	0	3675 (2/year)	2600 (1/year)	0	800 ^b	14,791
Interferon beta- 1b (Betaferon)	7716 (2/year)	0	0	0	3675 (2/year)	2600 (1/year)	0	800 ^b	14,791
Interferon beta- 1b (Extavia)	7716 (2/year)	0	0	0	3675 (2/year)	2600 (1/year)	0	800 ^b	14,791
Natalizumab (Tysabri)	0	16,250 (13/year)	0	0	3675 (2/year)	2600 (1/year)	0	5200 ª	27,725
Peg-interferon beta-1a (Plegridy)	7716 (2/year)	0	0	0	3675 (2/year)	2600 (1/year)	0	800 ^b	14,791
Teriflunomide (Aubagio)	0	0	0	0	3675 (2/year)	2600 (1/year)	448 °	800 ^b	7523

^a Analyses, MR, medical consultations and infusions will be done at the same day.

 $^{\rm b}$  Analyses, MR, and medical consultations will be done at the same day (2/year).

^c Every other month (numbers of medical consultations were deducted)

Drug	NAB-analyses	Infusion costs	Eye examina- tions	Startup costs	Medical consulta- tions	MRI	Blood tests (outpatient visits)	Travel costs	Total
Alemtuzumab ª (Lemtrada)	0	0	0	0	3675 (2/year)	2600 (1/year)	1232 (11/year; only for 35. year)	800 ^b	8307 (3 5.year) 7075 (+5.year)
Dimethyl fumarate (Tecifidera)	0	0	0	0	3675 (2/year)	2600 (1/year)	0	800 b	7075
Fingolimod (Gilenya)	0	0	0	0	3675 (2/year)	2600 (1/year)	0	800 b	7075
Glatiramer acetate (Copaxone)	0	0	0	0	3675 (2/year)	2600 (1/year)	0	800 b	7075
Interferon beta- 1a (Avonex)	0	0	0	0	3675 (2/year)	2600 (1/year)	0	800 ^b	7075
Interferon beta- 1a 44 mcg (Rebif)	0	0	0	0	3675 (2/year)	2600 (1/year)	0	800 ^b	7075

8.3: Monitoring costs associated with each of the treatments (beyond 2. year)

Interferon beta- 1a 22 mcg (Rebif)	0	0	0	0	3675 (2/year)	2600 (1/year)	0	800 ^b	7075
Interferon beta- 1b (Betaferon)	0	0	0	0	3675 (2/year)	2600 (1/year)	0	800 ^b	7075
Interferon beta- 1b (Extavia)	0	0	0	0	3675 (2/year)	2600 (1/year)	0	800 ^b	7075
Natalizumab (Tysabri)	0	16,250 (13/year)	0	0	3675 (2/year)	2600 (1/year)	0	5200 °	27,725
Peg-interferon beta-1a (Plegridy)	0	0	0	0	3675 (2/year)	2600 (1/year)	0	800 ^b	7075
Teriflunomide (Aubagio)	0	0	0	0	3675 (2/year)	2600 (1/year)	448 d	800 ^b	7523

^a The majority of patients receiving Alemtuzumab would not need new treatment after 5 –year treatment. It was assumed that 20% of patients need extra treatment (12 mg/day for 3 days) (expert opinion).

^b Analyses, MR, medical consultations and infusions will be done at the same day.

^c Analyses, MR, and medical consultations will be done at the same day (2/year).

^d Every other month (numbers of medical consultations were deducted)

## 9.1: The results of sensitivity analysis regarding stopping rule at EDSS=7 (discounted)*

	Total costs	Effects	Versus Interfe	Versus Interferon beta-1b 250 mg (Extavia)				
Drugs	(NOK)	(QALYs)	Incremental cost (NOK)	Incremental effect (QALYs)	ICER (NOK/QALY)	ICER (NOK/QALY)		
Interferon beta-1b (Extavia)	6,026,196	7.45						
Peg-interferon beta-1a (Plegridy)	6,290,635	7.64	264,439	0.19	1,424,765	1,424,765		
Natalizumab (Tysabri)	6,956,053	7.71	92,857	0.26	3,549,122	8,710,280		
Dominated therapi	es							
Interferon beta-1b (Beta- feron)	6,083,022	7.45	56,826	-	Dominated by interferon beta-1b (Ex- tavia)	Dominated by interferon beta- 1b (Extavia)		
Glatiramer ace- tate (Copaxone)	6,252,584	7.35	226,388	-0.10	Dominated	Dominated by interferon beta- 1b (Extavia) and interferon beta- 1b (Betaferon)		
Teriflunomide (Aubagio)	6,332,238	7.42	306,042	-0.03	Dominated	Dominated by interferon beta- 1b (Extavia), in- terferon beta-1b (Betaferon) and peg-interferon beta-1a		
Interferon beta-1a 22 mcg (Rebif)	6,500,898	7.24	474,702	-0.21	Dominated	Dominated by interferon beta- 1b (Extavia), in- terferon beta-1b (Betaferon), peg-		

						interferon beta- 1a, glatiramer acetate and teri- flunomide
Interferon beta- 1a 30 mcg (Avonex)	6,542,166	7.3	515,970	-0.15	Dominated	Dominated by interferon beta- 1b (Extavia), in- terferon beta-1b (Betaferon), peg- interferon beta- 1a, glatiramer acetate and teri- flunomide
Interferon beta-1a 44 mcg (Rebif)	6,572,277	7.36	546,081	-0.09	Dominated	Dominated by interferon beta- 1b (Extavia), in- terferon beta-1b (Betaferon), peg-interferon beta-1a, glati- ramer acetate and teriflunomide
Dimethyl fumarate (Tecifidera)	6,692,516	7.58	666,319	0.13	4,953,711	Dominated by peg-interferon beta-1a
Fingolimod (Gilenya)	7,034,538	7.47	1,008,342	0.03	40,301,928	Dominated by peg-interferon beta-1a, dimethyl fumarate and na- talizumab

* Alemtuzumab still was more effective (QALYS: 8.22) and less costly (Costs: 4,828,145) relative to other treatments (dominant strategy).

## 9.2: The results of sensitivity analysis using a 30-year time horizon of analysis (discounted) *

Drugs	Total costs	Effects	Versus Interferon beta-1b 250 mg (Extavia)			Sequential
	(NOK)	(QALYs)	Incremental cost (NOK)	Incremental effect (QALYs)	ICER (NOK/QALY)	ICER (NOK/QALY)
Interferon beta-1b (Extavia)	8,026,896	8.29				
Peg-interferon beta-1a (Plegridy)	8,276,892	8.55	249,995	0.26	960,134	960,134
Natalizumab (Tysabri)	9,033,436	8.64	1,006,540	0.36	2,818796	7,823,148
Dominated therapie	S	•		•		
Interferon beta-1b (Beta- feron)	8,090,003	8.29	255,409	-	Dominated by interferon beta-1b (Extavia)	Dominated by interferon beta-1b (Exta- via)
Glatiramer ace- tate (Copaxone)	8,282,305	8.17	331,284	-0.11	Dominated	Dominated by interferon beta-1b (Exta- via), interferon beta-1b (Beta- feron) and peg-interferon beta-1a
Teriflunomide (Aubagio)	8,358,180	8.26	565,095	-0.02	Dominated	Dominated by interferon beta-1b (Exta- via), interferon beta-1b (Beta- feron) and peg-interferon beta-1a
Interferon beta-1a 22 mcg (Rebif)	8,591,992	8.02	576,680	-0.327	Dominated	Dominated by interferon beta-1b (Exta- via), interferon

		-		r		
						beta-1b (Beta-
						feron), peg-in-
						terferon beta-
						1a, glatiramer
						acetate and
						teriflunomide
						Dominated by
						interferon
						beta-1b (Exta-
						via), interferon
Interferon beta-1a	8,603,576	8.12	611,852	-0.16	Dominated	beta-1b (Beta-
30 mg (Avonex)	0,003,570	0.12	011,052	-0.10	Dominated	feron), peg-in-
						terferon beta-
						1a, glatiramer
						acetate and
						teriflunomide
						Dominated by
						interferon
						beta-1b (Exta-
Interferon						via), interferon
beta-1a 44 mcg	9 639 7/9	8.19	717,167	-0.10	Dominated	beta-1b (Beta-
(Rebif)	8,638,748	0.19	717,107		Dominated	feron), peg-in-
(Rebil)						terferon beta-
						1a, glatiramer
						acetate and
						teriflunomide
Dimethyl fumarate						Dominated
(Tecifidera)	8,744,063	8.48	255,409	0.19	3,690,151	peg-interferon
(Technolera)						beta-1a
						Dominated by
						peg-interferon
Fingolimod (Gilenya)	9162,932	8.32	1,136,036	0.03	37,196,628	beta-1a, dime-
	9102,932	8.32		0.05		thyl fumarate
						and natali-
OALV: quality-adjus						zumab

 $\ast$  Alemtuzumab still was more effective (QALYS: 9.24) and less costly (Costs: 6,541,067) relative to other treatments (dominant strategy).

#### 9.3: The results of sensitivity analysis regarding "no EDSS improvement" (discounted) *

Drugs	Total costs	Effects	Versus Interferon beta-1b 250 mg (Extavia)			Sequential
	(NOK)	(QALYs)	Incremental cost (NOK)	Incremental effect (QALYs)	ICER (NOK/QALY)	ICER (NOK/QALY)
Interferon beta-1b (Extavia)	6,902,178	6.57				
Peg-interferon beta-1a (Plegridy)	7,109,166	6.73	206,988	0.16	1,309,477	1,309,477
Natalizumab (Tysabri)	7,706,752	6.78	804,573	0.20	3,935,743	12,890,581
		Dom	ninated therapies			
Interferon beta-1b (Beta- feron)	6,951,138	6.57	48,960	-	Dominated by interferon beta-1b (Extavia)	Dominated by interferon beta-1b (Exta- via)
Glatiramer acetate (Copaxone)	7,104,889	6.49	202,710	-0.08	Dominated	Dominated by interferon beta-1b (Exta- via), interferon beta-1b (Beta- feron) and peg-interferon beta-1a
Teriflunomide (Aubagio)	7,176,103	6.54	273,925	-0.03	Dominated	Dominated by interferon beta-1b (Exta- via), interferon beta-1b (Beta- feron) and peg-interferon beta-1a
Interferon beta-1a 22 mcg (Rebif)	7,329,592	6.4	427,413	-0.17	Dominated	Dominated by interferon beta-1b (Exta- via), interferon

		-				
						beta-1b (Beta-
						feron), peg-in-
						terferon beta-
						1a, glatiramer
						acetate and
						teriflunomide
						Dominated by
						interferon
						beta-1b (Exta-
						via), interferon
Interferon beta-1a	7 262 604	6.45	460 405	-0.12	Dominated	beta-1b (Beta-
30 mg (Avonex)	7,362,604	0.40	460,425	-0.12	Dominated	feron), peg-in-
						terferon beta-
						1a, glatiramer
						acetate and
						teriflunomide
						Dominated by
			477,998	-0.07	Dominated	interferon
						beta-1b (Exta-
Interferon						via), interferon
beta-1a 44 mcg	7,380,177	6,5				beta-1b (Beta-
(Rebif)						feron), peg-in-
						terferon beta-
						1a, glatiramer
						acetate and
						teriflunomide
Dimethyl fumarate					5,111,539	Dominated
(Tecifidera)	7,470,947	6.68	568,769	0.11	0,111,000	peg-interferon
(recilidera)						beta-1a
Fingolimod (Gilenya)				5 0.03 28,491,096		Dominated by
			865,925		28,491,096	peg-interferon
	7,768,104	6.60				beta-1a, dime-
		6.60		0.00		thyl fumarate
						and natali-
OALV: quality-adjus						zumab

 *  Alemtuzumab still was more effective (QALYS: 7.18) and less costly (Costs: 5,820,891) relative to other treatments (dominant strategy).

# 9.4: The results of sensitivity analysis regarding utility values (discounted) *

Drugs	Total costs	Effects	Versus Interferon beta-1b 250 mg (Exta		mg (Extavia)	Sequential	
	(NOK)	(QALYs)	Incremental cost (NOK)	Incremental effect (QALYs)	ICER (NOK/QALY)	ICER (NOK/QALY)	
Interferon beta-1b (Extavia)	6,035,711	7.88					
Peg-interferon beta-1a (Plegridy)	6,324,629	8.02	288,918	0.15	1,967,737	1,967,737	
Natalizumab (Tysabri)	7,000,849	8.08	965,138	0.21	4,649,607	11,131,827	
Dominated therapies	S						
Interferon beta-1b (Beta- feron)	6,094,252	7.88	58,541	-	Dominated by interferon beta-1b (Extavia)	Dominated by interferon beta-1b (Exta- via)	
Glatiramer acetate (Copaxone)	6,259,628	7.79	223,917	-0.08	Dominated	Dominated by interferon beta-1b (Exta- via), interferon beta-1b (Beta- feron) and peg-interferon beta-1a	
Teriflunomide (Aubagio)	6,353,620	7.84	317,909	-0.03	Dominated	Dominated by interferon beta-1b (Exta- via), interferon beta-1b (Beta- feron) and peg-interferon beta-1a	
Interferon beta-1a 22 mcg (Rebif)	6,511,148	7.69	475,437	-0.19	Dominated	Dominated by interferon beta-1b (Exta- via), interferon	

· · · · · · · · · · · · · · · · · · ·		1				
						beta-1b (Beta-
						feron), peg-in-
						terferon beta-
						1a, glatiramer
						acetate and
						teriflunomide
						Dominated by
						interferon
						beta-1b (Exta-
						via), interferon
Interferon beta-1a	6,556,702	7.74	520,991	-0.13	Dominated	beta-1b (Beta-
30 mg (Avonex)	0,550,702	1.14	520,991	-0.15	Dominated	feron), peg-in-
						terferon beta-
						1a, glatiramer
						acetate and
						teriflunomide
						Dominated by
						interferon
						beta-1b (Exta-
Interferon						via), interferon
beta-1a 44 mcg	6,586,671	7.79	550,959	-0.08	Dominated	beta-1b (Beta-
(Rebif)	0,000,071	1.19	550,959	-0.00	Dominated	feron), peg-in-
(Rebil)						terferon beta-
						1a, glatiramer
						acetate and
						teriflunomide
Dimethyl fumarate			679,345		6,182,526	Dominated
(Tecifidera)	6,715,056	7.99		0.11		peg-interferon
(recinicera)						beta-1a
Fingolimod (Gilenya)						Dominated by
			1,024,267	0.01	78,665,232	peg-interferon
	7,059,978	7.89				beta-1a, dime-
		7.89		0.01		thyl fumarate
						and natali-
OALV: quality adjust						zumab

* Alemtuzumab still was more effective (QALYS: 8.46) and less costly (Costs: 4,985,254) relative to other treatments (dominant strategy).

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