

Chapter 8. The NCBI BookShelf: Searchable Biomedical Books

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Summary

The [BookShelf](#) is a collection of biomedical books that can be searched directly in Entrez or found via keyword links in PubMed abstracts. Books have been added to the BookShelf in collaboration with authors and publishers, and the complete content (including all figures and tables) is free to use for anyone with an Internet connection.

The online books are displayed one section at a time, with navigation provided to other parts of the current chapter or to other chapters within the book. Many of the books on the BookShelf can be browsed without any restriction at all; others have less flexibility for navigating the complete content. The publisher (or the owner of the content) defines the rules for access.

The books are linked to PubMed through research papers citations within the text. In the future, more links may be established between the BookShelf and other resources at NCBI, such as gene and protein sequences, genomes, and macromolecular structures.

Content Acquisition

Basis for Inclusion

The BookShelf provides a venue through which publishers and authors can make the full text of biomedical books available to the scientific community. As the BookShelf grows, we welcome proposals from authors, editors, or publishers of any "in-scope" texts, from undergraduate textbooks to more specialized publications, including collections of review articles or workshop proceedings. The scope of the BookShelf is broadly biomedical, including clinical works and those concerning basic biological and chemical sciences.

Information for Authors, Editors, and Publishers

All books made available on the BookShelf have been provided in electronic format to NCBI. The publisher must provide the content *in full*. Because the complete content is

required for display and indexing purposes, the authors, editors, and publisher of a book should all be a part of the decision to participate in the BookShelf. Interested parties can contact us at books@ncbi.nlm.nih.gov for more information or to make a proposal for the inclusion of a book. There is a simple contract that specifies the terms of use of the content. A sample contract can also be obtained by request at the above email address.

The complete contents of each book will be converted into [XML](#) according to the NCBI Book Document Type Definition ([DTD](#)), a public domain DTD developed at NCBI for this project (see below). Books may be submitted to conform to this DTD, or NCBI will convert the source data to validate against the Book DTD. If a conversion needs to be done, the content must be in a format robust enough to meet the needs of the BookShelf publishing system and DTD.

Any book that was printed from [SGML](#) or XML should allow for a straightforward conversion. We have had success converting books from Word, XYWrite, PDF, and Quark Express formats, and we anticipate that we would also be able to convert from other desktop publishing packages. [HTML](#) and PDF formats are less desirable because the data formats are less detailed.

Figures should be supplied in TIFF format, although GIF and JPEG formats may be accepted. The submitted text files are converted into XML according to the NCBI Book DTD; graphic files are converted into GIF and JPEG formats. Three hard copies of the book are also required, along with the electronic files.

The XML files are stored in a database. When a reader requests a book, chapter, or section, the XML is retrieved from the database and converted into HTML on the fly using Extensible Stylesheet Language Transformations ([XSLT](#)) and Cascading Style Sheets ([CSS](#)).

How to Use the Books

There are three ways to access the content in BookShelf:

1. Through hyperlinked terms in PubMed abstracts
2. By a direct search using search terms or phrases (in the same way as the bibliographic database of PubMed is searched)
3. Through the Table of Contents of the book (note: some publishers restrict browsing through the entire book by disabling hyperlinks in the Table of Contents)

Links from PubMed

The BookShelf can be accessed from all PubMed abstract pages. When viewing a full PubMed abstract, select the **Books** hyperlink in the upper right-hand corner. This generates a version of the abstract in which certain phrases and terms appear as hypertext links (see Figure 1). The linked term may be one or more words in length. If a word or a

□ I: J Cell Biochem 2002;86(3):440-50 Related Articles, Links

Differentiation-dependent induction of CYP1A1 in cultured [rat small intestinal epithelial cells](#), colonocytes, and [human colon carcinoma cells](#): [Basement membrane-mediated apoptosis](#).

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[Rat small intestinal epithelial cells](#) and [human colon adenocarcinoma](#) cells cultured on Matrigel expressed the differentiation specific enzyme, sucrase-isomaltase, as determined by [indirect immunofluorescence](#). [Rat small intestinal epithelial cells](#), [rat](#) colonocytes, and [human colon adenocarcinoma](#) cells developed an altered morphology when cultured on Matrigel and became [apoptotic](#) within 24–48 h. [Benzo\[a\]pyrene](#) and 2,3,7,8-tetrachlorodibenzo-p-dioxin caused a 2- and 5-fold induction, respectively, of ethoxyresorufin-o-deethylase activity in [rat small intestinal epithelial cells](#) cultured on Matrigel. [Benzo\[a\]pyrene](#)- or 2,3,7,8-tetrachlorodibenzo-p-dioxin-induced ethoxyresorufin-o-deethylase activity in [rat small intestinal epithelial cells](#) cultured on plastic was not detected. 2,3,7,8-tetrachlorodibenzo-p-dioxin treatment caused a 14-fold induction of transfected, [rat CYP1A1-promoter-luciferase](#) activity in [rat small intestinal epithelial cells](#) cultured on Matrigel. [Benzo\[a\]pyrene](#) and 2,3,7,8-

Figure 1. A PubMed abstract showing terms linked to books. This view was generated by selecting **Links** found in the *top right corner* of the abstract and then selecting **Books** from the drop-down menu that appears.

phrase is linked, it means that the exact phrase also appears in at least one book. Selecting a linked phrase retrieves a list of books that contain that term.

A statistical weighting system based on the frequency of each phrase in a book section, relative to the rest of the book, is used to identify “good” phrases. A phrase that appears repeatedly in only a few sections and rarely in other parts of the book indicates a definitive phrase for those few sections; therefore, it ranks highly. Furthermore, the appearance of a phrase in the title, for example, has a greater value in the weighting system than one appearing solely in the text.

Each PubMed abstract can thus be linked to the appropriate book pages. This method allows two very dissimilar types of text—the dense, focused PubMed abstracts and the more descriptive book text—to find common ground.

Direct Search of Books

Book contents may be searched directly from the [BookShelf homepage](#) by using the search boxes located in the Table of Contents and navigation bars of books or by selecting **Books** from the pull-down menu in any Entrez database search bar (see [Chapter 15](#)). Search terms may be combined using [Boolean](#) operators that conform to PubMed syntax

(see [Chapter 2](#)). The BookShelf also allows search fields to be specified. A complete list of BookShelf database fields can be found in Table 1.

Table 1. Field limits for use in the BookShelf.

Field ^a	Use
[Author]	Search for the authors of books or chapters.
[Book]	Typically used with a Boolean expression to limit a search to a particular book.
[PmId]	Locate a journal article citation in a book by its PubMed ID.
[Rid]	Locate a particular book element (such as a figure or table) by its reference ID.
[Secondary Text]	Search for secondary text, e.g., units (mg/l, etc.)
[Title]	Search for words used in any title (book, chapter, section, subsection, figure, etc.).
[Type]	Locate a division of a book such as a section, chapter, or figure group.

^a Filters are applied immediately following a search term, with no separating spaces, e.g., watson[author] AND cmed[book].

Interpreting Results from a Search or PubMed Link

Results are shown as a list of books in which the term is found, along with the number of sections, figures, and tables that contain the term (Figure 2). The book that contains the most hits appears at the top of the list. Choosing the hyperlinked number of items that is associated with a particular book will then display a document summary list of the individual sections, figures, and tables found. (When a term or phrase is found fewer than 20 times within the BookShelf, the document summary page is shown directly, without first displaying the results clustered by book.)

The document summary list is sorted with the most relevant documents shown at the top of the page. The sorting makes use of scores allocated to phrases as a measure of how relevant they are to a given section (a part of the statistical weighting system also used for linking PubMed abstracts to the books). For each book section found, the title, along with some context regarding the hierarchy of the section location (e.g., the chapter and book), is given. An icon is used to distinguish figure legends and tables from text sections (Figure 3). Selecting a hyperlinked section title displays the part of the book that contains the search term. From this point, the user may be able to navigate further throughout the book content, according to the policy of the publisher (see *Navigating Book Content*).

Navigating Book Content

Each HTML page of content seen in a Web browser represents one section of one chapter of a book, i.e., all of the content (including subsections and so on) within the first-level heading of a chapter. The amount of content this represents varies according to the

NCBI **Bookshelf**

bMed Nucleotide Protein Genome Structure PopSet

Books dna Go Clear

Limits Preview/Index History Clipboard Details

Display Books Save Text Clip Add

709 items in **Molecular Cell Biology**. 4th ed.
Lodish, Harvey; Berk, Arnold; Zipursky, S. Lawrence; Matsudaira, Paul; Baltimore, David; ...
New York: **W H Freeman & Co**; c2000.

699 items in **Molecular Biology of the Cell**. 3rd ed.
Alberts, Bruce; Bray, Dennis; Lewis, Julian; Raff, Martin; Roberts, Keith; Watson, James D.
New York and London: **Garland Publishing**; c1994

616 items in **Cancer Medicine**. 5th ed.
Bast, Robert C.; Kufe, Donald W.; Pollock, Raphael E.; Weichselbaum, Ralph R.; Holland, J
Canada: **BC Decker Inc**; c2000

559 items in **Introduction to Genetic Analysis**. 7th ed.
Griffiths, Anthony J.F.; Miller, Jeffrey H.; Suzuki, David T.; Lewontin, Richard C.; Gelbart, V
New York: **W H Freeman & Co**; c1999.

426 items in **Modern Genetic Analysis**.
Griffiths, Anthony J.F.; Gelbart, William M.; Miller, Jeffrey H.; Lewontin, Richard C.
New York: **W H Freeman & Co**; c1999.

Figure 2. A results page from a BookShelf search. If more than 20 sections, tables, and/or figures are found that contain the query term, a summary page, such as the one above, is displayed.

structure of the original book. Some books have very long sections, some short, some a mixture; although on the whole, most chapters are divided into 3-10 sections.

The top of every page contains links to both short and detailed Tables of Contents and a description of the current location within the book (Figure 4). The hierarchal elements that describe the current location are hyperlinked and may be used to travel up the organizational levels of a book. Additionally, a navigation sidebar shows the current section among its peers and lists the figures and tables found within the current section (Figure 4). Reference citations in the text are linked where possible to PubMed abstracts by the [Citation Matcher](#). References internal to the book, e.g., to other chapters or

NCBI Bookshelf

bMed Nucleotide Protein Genome Structure PopSet

Books [dna AND cmed[book]] Go Clear

Limits Preview/Index History Clipboard Details

Display Summary Show: 20 Send to Text

Items 1-20 of 529 Page 1 of 27 Next

- 1: [Pulsed-Field Gel Electrophoresis](#)
Cancer Medicine e.5 -> Section 1. Cancer Biology -> 1. Molecular Biology -> Gene Analysis: DNA
- 2: [Intracellular Activation](#)
Cancer Medicine e.5 -> Section 13. Principles of Chemotherapy -> 44. Pharmacology-> General Mechanisms of Drug Action
- 3: [Mechanisms of Action](#)
Cancer Medicine e.5 -> Section 14. Chemotherapeutic Agents -> 47. Pyrimidine and Purine Antimetabolites -> Pyrimidine Analogues
- 4: [DNA sequencing using the chain...](#)
Cancer Medicine e.5 -> Section 1. Cancer Biology -> 1. Molecular Biology -> Gene Analysis: DNA
- 5: [Retroviral Vectors](#)
Cancer Medicine e.5 -> Section 3. Cancer Etiology -> 18. Tumor Viruses

Figure 3. A document summary list of book sections. The most relevant sections to the query appear at the top of the list. Note the icons that designate figure and table hits. The list may also be displayed in a brief format that lists only the section names that contain the term by choosing **Brief** from the drop-down menu to the immediate *right* of the **Display** button.

sections, figures, tables, and boxes, are also hyperlinked. Further navigation from the current page to other parts of the book depends on the access policy of the publisher.

Technology

Text Conversion and XML

All book content submitted to the BookShelf is converted to XML according to the public NCBI Book DTD.

Navigation

[About this book](#)

Section 2. Tumor Immunology

[10. Tumor Immunology](#)

[Targets for Immunotherapy](#)

[Issues](#)

[Problem Areas in Immunotherapy](#)

[Cancer Vaccines](#)

[Antibody Therapy](#)

[Adoptive Transfer](#)

→ [Cytokines](#)

[Conclusion](#)

[Acknowledgments](#)

[References](#)

Search

This book All books

PubMed

Cancer Medicine e.5 → Section 2. Tumor Immunology → 10. Tumor Immunology

Cytokines

Exogenously supplied cytokines are principally considered to provide immune regulation and to maximize the induction, amplification, and/or effector properties of the desirable immune response in the microenvironment of the vaccination site or the host-pathogen encounter. Furthermore, immune suppression or anergy imposed by the tumor-bearing state may alter the quantity and repertoire of endogenously produced cytokines essential for efficient clonal differentiation, maturation, and expansion. Consequently, cytokine-based immune intervention may be important for circumventing inhibitory mechanisms that disengage the development of a therapeutically relevant T-cell response. In general, cytokines that act best on immune cells act subsequent to antigen recognition and cellular activation, since those processes intimately serve to coordinate and govern cytokine receptor expression. On the basis of the spectrum and functional properties of distinct cytokines, IL-2, IL-12, IFN-g, and GM-CSF have been associated with cell-mediated immunity and tumor regression, while IL-4, IL-5, and IL-10 have been suggested to favor or promote tumor progression,^{46,47,108} which has been observed in both preclinical models and in carcinoma patients at different stages of disease progression.

Exogenous cytokines have been used as purified or recombinant proteins or expressed in recombinant viral vectors in conjunction with a given tumor antigen. Moreover, specific cytokine genes may be introduced directly into tumor cells, either stably, by transfection or transduction molecular techniques, or transiently, by infection with recombinant viruses. It is important to point out that the parameters employed for the use of exogenous cytokines in immunotherapy applications, such as dose, schedule, and frequency, are very complex and must be customized on the basis of their specific mode of action, pharmacokinetics, and predicted time of utilization during the course of the developing immune response.

IL-2 and IL-12 have been shown to play important roles in CD4⁺ Th1 and CD8⁺ CTL development and proliferation.^{75,77,108,180,200} Moreover, in experimental cancer models of adoptive or active immunotherapy, exogenous IL-2^{124,291,292} or IL-12^{181,293,294} has been shown to potentiate antitumor effects *in vivo*; these effects correlate with enhanced CTL activity. GM-CSF has also been reported to enhance antigen-specific T-cell responses, such as proliferative, CTL, and delayed-type hypersensitivity reactions^{141,295} and antitumor responses.^{67,87,206-208} It should be noted, however, that GM-CSF most likely acts indirectly via the recruitment and activation of host APC populations, such as macrophages and DCs.^{3,6,63} Increased APC

Figure 4. Navigation elements that appear on every book HTML page. Hyperlinks to various sections within a chapter appear within a navigation bar to the *left* of each page. Hyperlinks may be disabled within some books at the request of the publisher. A **Search** box is located *below* the navigation bar. At the *top* of each page is a hyperlinked, hierarchical tree that illustrates a page's relative position within a book.

Any files submitted in SGML or XML are converted to the Book DTD using **XSLT**. Books submitted in desktop publishing or word processing formats are converted by a contractor using proprietary technologies. Once the book XML is valid against the DTD, the book is ready to be loaded into the database.

The XML for a book is generally a set of files. Each chapter and appendix is an independent file, as is the frontmatter. The book is pulled together by the book.xml file, which defines the structure of the book. For example, a book with two chapters and a bibliography at the end of the book would be structured as follows:

```
<!DOCTYPE book SYSTEM "ncbi-book.dtd" [
<!-- graphics -->
```



```

<!ENTITY % Graphics SYSTEM "graphics.xml">
%Graphics;
<!ENTITY frontmatter SYSTEM "fm.xml">
<!ENTITY chapter1 SYSTEM "ch1.xml">
<!ENTITY chapter2 SYSTEM "ch2.xml">
<!--back-->
<!ENTITY biblist SYSTEM "biblist.xml">
]>
<book>
&frontmatter;
<body>
&chapter1;
&chapter2;
</body>
<back>
&biblist;
</back>
</book>

```

The book.xml file is composed of two parts. The first part defines all of the components that are required to build the book. These definitions occur within the <!DOCTYPE [] > tag. The second part builds the structure of the book. The root element is <book>. <book> contains whatever is in &frontmatter;, <body>, and <back>.

The book.xml example above refers to five external files: fm.xml, which contains all of the frontmatter of the book; ch1.xml, which contains chapter 1; ch2.xml, which contains chapter 2; biblist.xml, which contains the bibliography for the book; and graphics.xml, which defines the images. If any of these files is not valid according to the DTD or if the files are not found where they are defined in their <!ENTITY > declaration, then the book will not be valid.

Images

All of the images, including figures, icons, and book-specific character graphics (see Special Characters below), are called out in the text as entities. The entities are defined in the graphics.xml file.

```

<!ENTITY ch2fu6 SYSTEM "data/mga/pictures/ch2/ch2fu6.gif" NDATA GIF>
<!ENTITY ch2fu7 SYSTEM "data/mga/pictures/ch2/ch2fu7.jpg" NDATA JPG>
<!ENTITY ch2fu8 SYSTEM "data/mga/pictures/ch2/ch2fu8.gif" NDATA GIF>
<!ENTITY ch2fu9 SYSTEM "data/mga/pictures/ch2/ch2fu9.gif" NDATA GIF>
<!ENTITY ch2fu10 SYSTEM "data/mga/pictures/ch2/ch2fu10.gif" NDATA GIF>
<!ENTITY ch2e1 SYSTEM "data/mga/pictures/ch2/ch2e1.gif" NDATA GIF>
<!ENTITY ch2e2 SYSTEM "data/mga/pictures/ch2/ch2e2.gif" NDATA GIF>

```


Graphic files are converted into GIF and JPEG formats and optimized for display on the Web. The images are not loaded into the database; they are retrieved from a file server when called by the HTML page.

Math and Formulae

Math expressions and chemical formulae and structures are handled as images.

Special Characters

The BookShelf uses the same character sets that PubMed Central uses (see [Chapter 9](#)). These include a number of standard ISO character sets (8879 and 9573), along with a set of characters that has been defined to accommodate characters not in the standard set. The ISO Standard Character sets referenced are listed in Box 1 of [Chapter 9](#). Special characters are converted to the BookShelf/PubMed Central (PMC) character set during conversion into XML. Characters created for one book (book-specific characters) are called out in the XML as images. To provide for the most flexibility in displaying characters across platforms, BookShelf uses UTF-8 encoding whenever possible. Because not all browsers support the same subset of UTF-8 characters and some characters cannot be represented in UTF-8, the BookShelf displays characters as a combination of GIFs and UTF-8 characters, depending on the Browser/OS combination and the character to be displayed.

The BookShelf Data Flow

The XML files for each book are stored in a Sybase database. To show the best representation to each reader, the reader's browser and operating system are noted and passed to the rendering software. When a reader requests a book, chapter, or section, the XML and the character images or UTF-8 characters appropriate to the reader's system are retrieved from the database.

The XML is converted to HTML using XSLT stylesheets. The look of the HTML pages is controlled further using CSS, which allow manipulation of colors, fonts, and typefaces (Figure 5).

The Table of Contents for a book is created from actual elements within the book content, rather than from the Table of Contents given in the book frontmatter of the hard copy. This ensures that the Table of Contents represents the content accurately as it is organized on BookShelf.

NCBI Book DTD

History

In the first version of the NCBI BookShelf project, Quark files were converted directly to HTML for display online. The result was effective, illustrating the value of having a

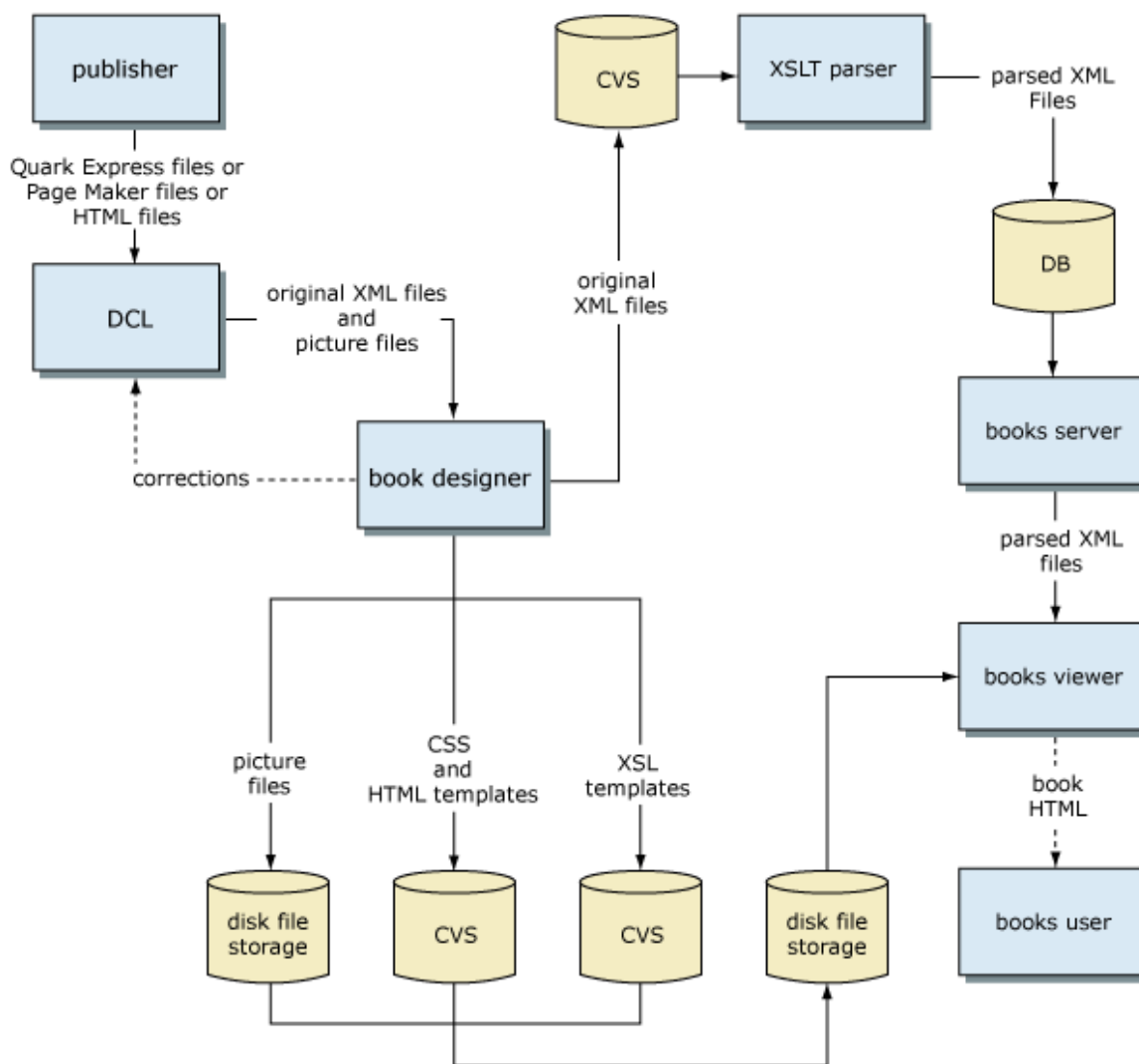


Figure 5. Processing and data flow of books.

textbook online and linked to PubMed; however, it was labor intensive, limiting, and not scaleable.

To simplify the delivery of books online and to allow for the expansion of linking within the Entrez system, NCBI decided to convert all content into a centralized XML format.

The normalized XML content is easier to render, allows added value such as the addition of links to other NCBI databases, and simplifies the addition of new volumes.

PMC created a new DTD for the BookShelf project, which was based on the ISO-12803 DTD. As more books were converted to the NCBI Book DTD, changes had to be made to accommodate the data.

The NCBI Book DTD is a public DTD available on request from books@ncbi.nlm.nih.gov.

Frequently Asked Questions

1. How do I access the books at NCBI?

The online books can be accessed by direct searching in Entrez or through PubMed abstracts. After performing a general PubMed search, click on the author name of one of the search results to view the abstract. A hypertext link called **Links** is displayed to the right of the abstract title. This link contains a drop-down menu consisting of various choices, depending upon the specific abstract. Choosing **Books** from this drop-down menu will highlight keywords in the abstract that, when selected, initiate a search of all BookShelf content for that particular term.

2. Which books are available at NCBI?

The book list is updated on a regular basis and can be viewed on the BookShelf homepage: <http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?db=Books>.

3. Can I search the books at NCBI?

Yes, the books can be searched either as a complete collection or as a single, selected book (restricted using search options found under **Limits**).

4. Can I browse the whole book?

The system has been designed so that the user is delivered to the most relevant book sections for a particular term or concept. Although navigation is possible in the immediate vicinity of the page to which you are delivered, it may not be possible to browse the complete book on BookShelf. The range of navigation for each book is determined on a case-by-case basis, in agreement with the publisher.

5. I am the publisher/author/editor of a book. How can I participate?

Please email books@ncbi.nlm.nih.gov to discuss potential projects.