


新版电子图书阅读器

Books@Ovid 推出的新书阅读器简洁易用，帮助用户方便高效的阅读并获取电子图书，增强的图书阅读器功能包括：


- 更新了阅读视图、搜索、电子书导航、个性化自定义等功能。
- 为各种设备提供优化的体验，会根据设备的屏幕进行显示调整。
- 现在，您可以直接从目录中的链接转到标题中的特定部分，如章节或附录。
- 您可以根据自己的喜好调整背景、文本颜色、字符大小、间距和屏幕上的字符对齐方式。

新图书阅读器主屏幕

 全屏显示

 章节导航


 图书目录


 图书检索

 图书引文

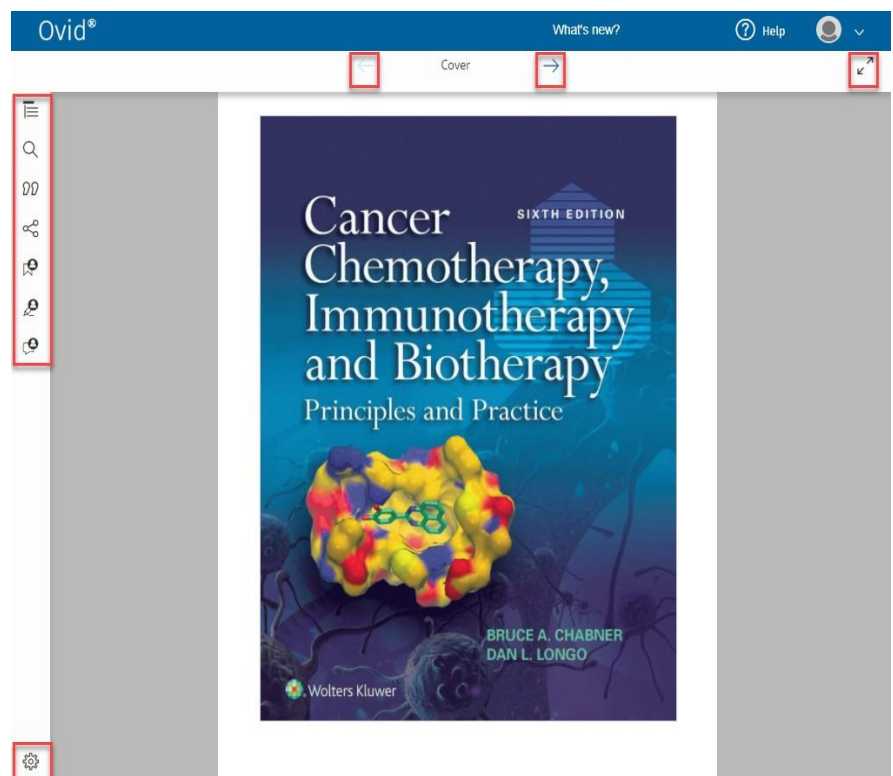
 内容分享

 书签

 要点标记

 注释

 页面设置



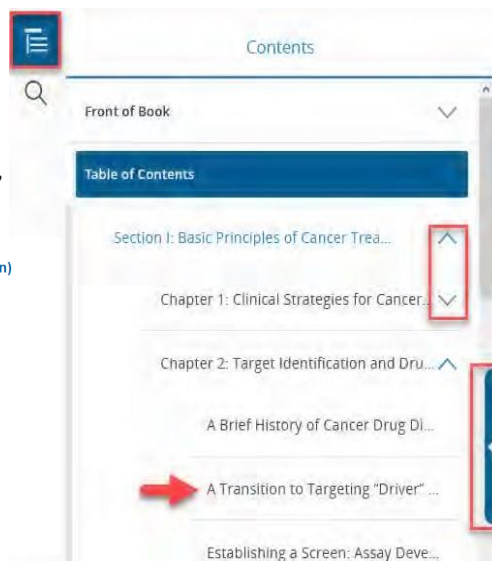
在 Ovid 平台上直接点击图书上的绿色背景图标，进入图书阅读器。




5-Minute Clinical Consult 2011, The (19th Edition)


Domino, Frank J.
Lippincott Williams & Wilkins, 2010
ISBN: 9781608312597, 1608312593
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新版图书阅读器



图书目录

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 隐藏目录

★ 选择具体章节，内容显示在主屏幕上。

内容视图:

many other oncogenic drivers. Specificity of inhibitors for specific oncogenic proteins has been a recurrent challenge, as many of the drugs in current clinical practice have multiple sites of action, leading both to additional antitumor uses, but also to toxicities (see Table 2.1). Since there is high homology of the ATP binding sites of subset of receptor tyrosine kinases (Fig. 2.6) toxicities to (Print pagebreak 24) (Print pagebreak 25) (Print pagebreak 26) skin and GI epithelium are common side effects for compounds of this class. However, it has been possible to synthesize drugs highly specific for a single, specific target unique to cancer cells, as for example osimertinib,²⁴ a highly active and specific inhibitor of the T790M mutant *EGFR* gene. It is 100-fold more potent against the mutant enzyme as compared to the wild-type *EGFR*.

- 滑动框
- 章节内容 PDF 下载
- 章节图片
- [\(Fig. 2.6\)](#) 图片或图标链接
- ★ 图片、图像、图表可以单击放大显示，并可进行图片或 PPT 文件下载。

图书检索:

imatinib binding

However, while **imatinib binding** (A) prevents the further conformational binding of a juxtamembrane (JM domain) into the pocket, the more

Disruption of **imatinib binding** by the T315I mutation. The bulky isoleucine side chain causes steric hindrance, preventing drug access

However, while **imatinib binding** (A) prevents the further conformational binding of a juxtamembrane (JM domain) into the pocket, the more

页面设置:

Reader Settings

Text size adjustment: Tr ——— Tr

Align: [Left] [Center] [Right] [Justify]

Font: Default

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Mode: [Tr] [Tr] [Tr]

Margins: [Default] [Wide] [Narrow]

Scroll View: OFF ON

Reset to default

- 文字大小调整
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- 滚动阅读关/开
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