Abstract
The equivalence of the US Orange Book in China, the “CN Orange Book” has been operating since 29 June 2021. This “CN Orange Book” is a mountain full of treasures, for example clear indication on what claim is covering which subject matter in a particular drug patent. This “unofficial” user guide aims to provide foreigners information on how to navigate this “CN Orange Book”, which operates in Chinese. This is the only tool available to a new drug owner to delay approval of the corresponding generic drugs marketing approval process, as China does not provide new drug/biologic exclusivity.

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Contents
“Unofficial” CN Orange Book user guide for pharmaceutical patents (June 2022 ver.) ....3
Available to anyone, but could only search in Chinese ........................................4
Extensive information contained therein – good for FTO research ..................................6
Restrictions on the type of patents that could be registered .............................................10
Only Chinese company could register, and make entries on the platform ......................11
Types of declaration from a generic drug applicant against an innovative drug ............11
The first type 4.2 declaration ..................................................................................14
Drug patent linkage complaint situation in China in Dec 2021 ....................................17
Current drug patent linkage complaint situation in China in June 2022 .......................18
  Beijing IP Court first drug patent linkage decision .................................................19
  CNIPA “first” drug patent linkage decision ..........................................................20
  Observations from CN drug patent linkage decisions as of June 2022 .....................21
Comparison between the US and CN drug patent linkage system ................................22
Conclusion ..............................................................................................................22
Appendix - Differences between the US and China drug patent linkage systems ........24
“Unofficial” CN Orange Book user guide for pharmaceutical patents (June 2022 ver.)

In the US, the Orange Book is a publication from the US FDA that identifies drug products with marketing approval from the FDA and related patent and exclusivity information. Information in the Orange Book could be used in actions under the Hatch-Waxman Act, which allows a patentee to file an artificial act of infringement that allows the drug innovator to file suit before the generic drug is commercialized, such that the drug marketing approval process of the generic drug at the US FDA could be suspended.

China is required to introduce a similar system under the Phase-I Trade Agreements signed with the US in January 2020. I have been following this since then, and have several articles and posts on this topic. Recently, this “CN Orange Book” (official name: China listed drugs patent information registration platform) has become officially online on 29 June 2021, with many developments thereafter, and I have received many questions on how to use this. This motivates me to write this “unofficial” user guide, mainly for foreign users. This user guide may need updates periodically as things are changing too fast in China.

This drug patent linkage system is the only tool available to a new drug owner to delay approval of the corresponding generic drugs marketing approval process, as China does not provide new drug exclusivity (5 years for chemical, and 12 years for biologic) as in the US.

The equivalence of the US FDA in China is the China National Medical Products Administration, the NMPA.

This is the June 2022 version of this guide, as there have been many developments regarding the first drug patent linkage decisions after my various versions below:

- first July 2021 version finished on 19 July 2021;
- second October 2021 version finished on 14 October 2021; and
- third December 2021 version finished on 5 December 2021.
Available to anyone, but could only search in Chinese
The “CN Orange Book” is available at https://zldj.cde.org.cn/home. Anyone can access information therein. Below is a screen shot of the first page showing different functions thereon:
For searching drug patent information already registered, the information is divided into three tabs, each respectively directed to Chinese medicine, chemical drug, and biological drugs as shown below (with the tab of chemical drugs chosen):

<table>
<thead>
<tr>
<th>序号</th>
<th>药品名称</th>
<th>规格</th>
<th>批准文号/注册证号</th>
<th>剂型</th>
<th>操作</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>国药准字20190024</td>
<td>化学药品</td>
<td>盐酸苯海索片0.5mg</td>
<td>片剂</td>
<td>查看</td>
</tr>
<tr>
<td>2</td>
<td>国药准字20190002</td>
<td>化学药品</td>
<td>碳酸钙片</td>
<td>片剂</td>
<td>查看</td>
</tr>
<tr>
<td>3</td>
<td>国药准字20190091</td>
<td>化学药品</td>
<td>氨酚烷胺</td>
<td>片剂</td>
<td>查看</td>
</tr>
<tr>
<td>4</td>
<td>国药准字20180077</td>
<td>化学药品</td>
<td>布洛芬片20mg</td>
<td>片剂</td>
<td>查看</td>
</tr>
<tr>
<td>5</td>
<td>国药准字20180076</td>
<td>化学药品</td>
<td>布洛芬片15mg</td>
<td>片剂</td>
<td>查看</td>
</tr>
<tr>
<td>6</td>
<td>国药准字20180075</td>
<td>化学药品</td>
<td>布洛芬片10mg</td>
<td>片剂</td>
<td>查看</td>
</tr>
<tr>
<td>7</td>
<td>国药准字20180021</td>
<td>化学药品</td>
<td>布洛芬片</td>
<td>片剂</td>
<td>查看</td>
</tr>
<tr>
<td>8</td>
<td>国药准字20180020</td>
<td>化学药品</td>
<td>布洛芬片</td>
<td>片剂</td>
<td>查看</td>
</tr>
<tr>
<td>9</td>
<td>国药准字20171054</td>
<td>化学药品</td>
<td>氨酚烷胺</td>
<td>片剂</td>
<td>查看</td>
</tr>
</tbody>
</table>

共468条
Searches could be done and directed to the drug name, the format of the drug, dosage, and the drug marketing number at the NMPA. Mostly the name of the drug is used. Partial search for drug patent information already registered is available, but could only be done in Chinese. For example, using only the last three words of Palbociclib in Chinese (“柏西利” in “哌柏西利”) resulted in the following search results:

**Extensive information contained therein – good for FTO research**

The “CN Orange Book” has the following number of entries as of different dates in the table below:

<table>
<thead>
<tr>
<th>Type of drugs</th>
<th>30 Jun 2021 no.</th>
<th>19 Jul 2021 no.</th>
<th>3 Oct 2021 no.</th>
<th>5 Dec 2021 no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chinese medicines</td>
<td>30</td>
<td>143</td>
<td>243</td>
<td>280</td>
</tr>
<tr>
<td>Chemical drugs</td>
<td>278</td>
<td>448</td>
<td>532</td>
<td>568</td>
</tr>
<tr>
<td>Biological drugs</td>
<td>39</td>
<td>60</td>
<td>78</td>
<td>80</td>
</tr>
</tbody>
</table>

There was a relatively big surge in July 2021, shortly after the database was firstly introduced on 29 June 2021, while the number of registrations slowed down significantly thereafter, particularly after October 2021. This could be expected, as every drug owner would rush into the register at the beginning.
One amazing feature of the “CN Orange Book” is the inclusion of extensive information in each entry. Below is a screenshot of the entry in the “CN Orange Book” for the drug Palbociclib capsule 125mg (Note: only part of the entries of one of the three patents registered for this drug is shown below due to limit of the screen shot that could be taken):

![Entry Screenshot]

It can be seen that in addition to the patent number, the following information in the “CN Orange Book” database entries is also available:

1) The granted patent itself is available for downloading at a link on the same page.
2) The claim number(s), and the relevant subject matter, are clearly identified.
3) The full-term expiry of the patent(s) is clearly identified.
4) Whether the patent has survived through invalidation is also specified in the note section.
On the other hand, there exists entries without the above extensive information of 1)-4), as shown below as of 19 July 2021:
The same entry above has its information updated as of 3 October 2021, indicating claims 5 to 8 of the relevant patent CN ZL03821144.0 are directed to the drug composition containing the active pharmaceutical ingredient API vardenafil hydrochloride trihydrate:

It should be noted that the relevant regulations, the NMPA Measures (see below), put the onus on the patentee/drug marketing approval holder DMAH to ensure the accuracy of the information registered on the platform, which requires the claim(s) covering what aspect of the drug to be identified (Article 4 of the NMPA Measures). Failure to do so could be legally liable (Article 15 of the NMPA Measures), while it is unclear what exactly are the punishments.

Most of the entries in the “CN Orange Book” contain extensive information like the one of Palbociclib capsule 125mg above (if not, updated could be expected). Such information could be very useful for a freedom-to-operate (FTO) exercise. The immediate availabilities of the grant patent publication and the full-term expiry are already very useful. The specific identifications of the claim(s) and relevant subject matter are even more so.
By contrast, the equivalent FDA records only contain the US patent numbers and the full-term expiry date, but nothing further. Below is the screenshot of Palbociclib capsule 125mg in the US FDA Orange Book records:

**Restrictions on the type of patents that could be registered**
The “CN Orange Book” restricts that only the following types of patents could be registered for the respective drugs (Articles 5 and 12 of the NMPA Measures):

- Chemical drugs – chemical compound of the active ingredient; composition comprising the active ingredient; medical use
- Biological drug – sequence structure of the active ingredient; medical use
- Chinese medicine – composition; extracts from Chinese medicine; medical use

As biologics are also included in the Chinese platform, such is in fact a combination of the US Orange (for chemicals) and Purple (for biologics) Books.

While not specified in the articles of the NMPA measures, the accompanying explanations of the NMPA measures specify that patents covering intermediates, metabolites, crystal form, method of manufacturing and testing methods are not considered as relevant patents under the patent linkage system, i.e. these patents should not be registered on the “CN Orange Book”. Therefore, the above types of patents would not be seen in the “CN Orange Book”.

Having said so, one way to get around the restriction on crystal form is to record a patent having a composition claim, or medical use claims (in the form of Swiss-type claim, as method of treatment is not patentable in China) containing the crystal form of the API. An example will be shown below in the first contentious drug patent linkage declaration.
Only Chinese company could register, and make entries on the platform
While everyone could access the platform to view information therein, in order to register entries on the platform (either for a drug, or making a declaration against a drug), one must register as a user on the platform. However, only a Chinese company could do so, as it is required to input the so-called "unified social credit codes" (compulsory) during registration, which are available to all Chinese companies. Further, the registration requires a CN mobile number to complete (for receiving a confirmatory text message).
See the screenshot below:

Types of declaration from a generic drug applicant against an innovative drug
For details on how such a declaration works, please refer to my articles below:

https://www.linkedin.com/pulse/cn-patent-inkage-measures-trial-finalized-issued-nmpa-toby-mak (This article talks about the finalized NMPA measures, which is based on the draft measures below)
https://www.linkedin.com/pulse/cn-drug-patent-registration-system-orange-book-opened-toby-mak (This article talks about the draft NMPA measures)
As a summary, when a generic drug applicant applies for drug marketing approval at the NMPA, the generic drug applicant is required to make a declaration of any one of the following four types:

1. That there is no patent information registered related to the drug.

2. That the patent(s) has already expired, or been declared invalid.

3. The date on which the patent(s) will expire, and that generic applicant promises that the generic drug will not go on the market until the expiry of the patent(s).

4. That the patent(s) is not infringed, or should be declared invalid.

As of 14 June 2022, 1095 declarations have been announced on the CN platform, showing a significant increase from 410 declarations on 3 October 2021 and 745 declarations on 5 December 2021. The vast majority of the declarations are types 1 to 3 declarations. The first type 3 declaration was announced on 9 July 2021, with the screen shot below:

![Screen shot of declaration type 3](image)

The key is the type of the declaration filed, in the screen shot above being type 3. It should also be noted that this page subdivides declaration type 4 into the following two subtypes:

- **4.1** - The patent(s) should be declared invalid.
- **4.2** – The generic drug does not fall within the scope of the patent(s).
The type of the declaration determines whether the drug patentee/DMAH could take action against the subject generic drug marketing approval application. It appears only action could be taken against declaration type 4.2 (the generic drug does not fall within the scope of the patent(s)). While Article 7 of the NMPA measures could be literally read as allowing complaints to be filed at a court or the China National Intellectual Property Administration (the CNIPA) for all 4 types of declarations, when combining with the relevant measures of the Supreme People’s Court and the CNIPA, it appears that there is a typographical error in Article 7 of the NMPA measures, and complaint could only be filed for declaration type 4.2.

Therefore, if a generic drug applicant filed a type 4.1 declaration, and even if the generic drug applicant did not file a subsequent invalidation petition, the drug marketing approval process would still proceed as normal by the NMPA as if drug patent linkage protection does not exist.

On the other hand, if the generic drug applicant did file a subsequent invalidation petition and could successfully invalidate the patent, a 12-months exclusivity period would be offered to the first successful challenger, i.e., drug marketing approval would not be granted to another generic drug applicant during this 12-months period.
Type 4.1 declaration has already been announced, with an example shown below:

In the above, the drug at issue is ticagrelor 90mg tablet, with AstraZeneca being the DMAH, and the generic drug applicant being Shanxi Deyuantang Pharmaceutical Co., Ltd. (Deyuantang). As mentioned above, in this situation, there is nothing AstraZeneca could do to delay the drug marketing approval process of the generic ticagrelor 90mg tablet from Deyuantang at the NMPA.

The first type 4.2 declaration
This was announced on 23 July 2021 directed to the drug dapagliflozin (达格列净) 5 and 10 mg, with AstraZeneca as the DMAH in China. The relevant screen shot for the 5mg drug is shown below, while that of the 10 mg drug has similar contents.
It is interesting to note that the generic drug applicant Jiangsu Hansoh Pharmaceutical Group Co., Ltd. (江苏豪森药业集团有限公司, Hansoh) filed a type 3 declaration against CN200910158686.6 directed to the chemical compound of the API dapagliflozin itself, i.e. Hansoh undertakes that they would only put their drug onto the market after CN200910158686.6 has expired on 15 May 2023 (CN200910158686.6 is a divisional from CN03811353.8).

On the other hand, Hansoh filed type 4.2 declarations against the following 3 CN patents, i.e. Hansoh declared that their generic drug does not fall within the scope of the following 3 CN patents that are directed to specific formulations and medical uses:

1) CN200880016902.7 (full term expiry 21 March 2028) - specific formulation comprising the API in the form of dapagliflozin propylene glycol hydrate, microcrystalline cellulose, lactose, crospovidone, silicon dioxide, magnesium stearate, with each components presents in respective specific amounts (claim 8).

2) CN200780024135.X (full term expiry 21 June 2027) - Medical use (Swiss-type claim) of dapagliflozin propylene glycol hydrate with specific crystalline structure (note: as mentioned above patents directed to crystalline structure of a drug per se should not be registered in the CN Orange Book) in treating diabetes, insulin resistance, hyperglycemia, hyperinsulinemia, elevated blood levels of fatty acids or glycerol,
hyperlipidemia, dyslipidemia, obesity, hypertriglyceridemia, or diabetic complications (claim 9).

3) CN201210201489.X (full term expiry 21 March 2028) - specific formulation comprising the API in the form of dapagliflozin propylene glycol hydrate with a specific dosage regime (claim 1), with further components including bulking agent, binder, disintegrant, and so on (claims 2 and 3), and the medical uses of such specific formulation (Swiss-type claims) for treating diabetes.

For 2), it should be noted that the patent is in fact directed to the crystal form of the API in the form of dapagliflozin propylene glycol hydrate, and therefore the CN Orange Book registration is directed to the medical use in Swiss-type claim containing such specific crystal form. Therefore, for China, it is important to include at least one composition or medical use Swiss-type claim for drug crystal form patent.

According to the above, Hansoh chose to challenge the patents directed to specific formulations comprising the API dapagliflozin propylene glycol hydrate, and the medical use of the crystalline API. Comparing with the chemical compound patent CN200910158686.6 (which Hansoh chose to deal with using a type 3 declaration), there is higher chance to get around these formulation and medical use patents, for example by not using dapagliflozin propylene glycol hydrate, but uses dapagliflozin itself (whether this would be caught by equivalence is another question). In fact, Hanosh declared that the API in their generic drug is different from that in these 3 CN patents, i.e. dapagliflozin propylene glycol hydrate.

AstraZeneca had until 6 September 2021 (45 days from 23 July 2021) to file a complaint against Hansoh's type 4.2 declaration against the 3 CN patents above at the Beijing IP Court, or at the CNIPA. Failure to do so would result in that the drug marketing approval application would be processed as normal by the China National Medical Products Administration as usual, i.e. without any delay.

If AstraZeneca filed the complaint and was accepted (by the Beijing IP Court, or the CNIPA), and a decision was made within 9 months from the date of acceptance of the complaint:
• if the decision was in favor of AstraZeneca, the drug marketing approval would be delayed to shortly before the expiry of the above three patents, i.e. maximum delay up to 2028.
• if the decision was not in favor of AstraZeneca or not within 9 months, the drug marketing approval would be processed as normal with no delay.

However, as Hanosh undertakes a type 3 declaration against CN200910158686.6, the earliest date that Hanosh could put their generic drug on the market in China is 15 May 2023.

**Drug patent linkage complaint situation in China in Dec 2021**
For the first type 4.2 declaration discussed above, I have tried to check whether AstraZeneca had filed the complaint at the Beijing IP Court the CNIPA by 6 September 2021, but no luck. In light of the relative ease of getting around the above three subject patents, and Hansoh undertakes that they would only put their drug onto the market after 15 May 2023, AstraZeneca may choose not to file the compliant against the type 4.2 declarations.

On the other hand, according to the announcements from the CNIPA and the Beijing IP Court at the respective links below, 12 drug patent linkage complaints (out of 23 filed) have been accepted by the CNIPA as of October 2021, while 1 has been accepted by the Beijing IP Court as of November 2021 (not clear how many were filed at the Beijing IP Court and were rejected).

- [https://mp.weixin.qq.com/s/0PNtsF0XBuwxULqYgnV-Uw](https://mp.weixin.qq.com/s/0PNtsF0XBuwxULqYgnV-Uw)

The CNIPA did not provide any details on the complaints they accepted, but the Beijing IP Court did. Specifically, the drug at issue of the complaint accepted by the Beijing IP Court is Eldecalcitol of the Japanese company Chugai Pharmaceutical Co., Ltd. (under the brand Edirol) for treating osteoporosis. The generic supplier is Wenzhou Haihe Pharmaceutical Industry. The patent at issue is CN ZL2005800098777.6.
Apparently the CNIPA has accepted more cases than the BJ IP Court. After checking with the CNIPA, the reason appears to be that the CNIPA does NOT require notarization and legalization of formality documents including the Power of Attorney and the business registration certificates to accept the complaints. Specifically, the CNIPA only requires Power of Attorney with a simple signature, while only notarization is required for the business registration certificates, to accept the complaint.

By contrast, the BJ IP Court requires notarization AND legalization of the above formality documents in order to accept the complaint. Practically, it is very difficult, if at all possible, to fulfil these requirements within 45 days, especially during the current pandemic.

Such discrepancies on handling of formality documents are reflected on the differences in the numbers of cases accepted (CNIPA 12, BJ IP Court 1).

If the CNIPA continues with this practice, such could make CNIPA a more preferable forum for filing drug patent linkage complaints than the Beijing IP Court. This is because a drug patent linkage complaint must be filed within 45 days from the date of announcement of the relevant 4.2 declaration, otherwise there would be no delay on the drug marketing approval process at the NMPA, i.e. as drug patent linkage protection does not exist. Therefore, the CNIPA is the only choice after the complaint is rejected by the BJ IP Court.

Assuming that the CNIPA made the announcement shortly after they accepted the first drug patent linkage complaint, if a decision was made on the first case by July 2022 and found the generic drug to fall within the scope of the patent, then the NMPA should stop the drug marketing approval process until 20 days before the patent expired. Otherwise, the drug marketing approval process would proceed as usual.

**Current drug patent linkage complaint situation in China in June 2022**

In April 2022, both Beijing IP Court and the CNIPA have announced that they have made their respective first drug patent linkage decision, with Beijing IP Court made the first announcement on 15 April 2022, and the CNIPA followed suit on 25 April 2022 at the links below (looks like the Beijing IP Court and the CNIPA were competing to announce):

- Beijing IP Court - [https://mp.weixin.qq.com/s/-9SocP2TLE20Xw7me3OxNw](https://mp.weixin.qq.com/s/-9SocP2TLE20Xw7me3OxNw)
Both decisions ruled in favor of the generic supplier. While there were critics that these are yet another examples that China is not protecting intellectual property properly, the patents relied on by the DMAH appear to be able to be get around relatively easily, resulting in these decisions. Details of these cases will be discussed below.

**Beijing IP Court first drug patent linkage decision**
Details of the case are as below:

- The drug at issue is Eldecalcitol of the Japanese company Chugai Pharmaceutical Co., Ltd. (under the brand Edirol) for treating osteoporosis.
- The generic supplier is Wenzhou Haihe Pharmaceutical Industry (Haihe).
- The patent at issue is CN ZL2005800098777.6 ('777).

As mentioned above, Haihe was found by the Court to be not falling within the scope of ‘777 literally or by equivalence, for which claim 1 specifies the following:

- The degradation of the active ingredient ((5Z,7E)-(1R,2R,3R)-2-(3-hydroxypropoxy)-9,10-secocholesta-5,7,10(19)-triene-1,3,25-triol) into specific degradation products (6E-(1R,2R,3R)-2-(3-hydroxypropoxy)-9,10-secocholesta-5(10),6,8(9)-triene-1,3,25-triol and/or (5E,7E)-(1R,2R,3R)-2-(3-hydroxypropoxy)-9,10-secocholesta-5,7,10(19)-triene-1,3,25-triol) is suppressed.
- After being stored for 12 months at room temperature and shading, the amount of the degradation product is 1% or less.

It may not be that difficult for the generic of Haihe to get around claim 1 of ‘777. For example, this could be achieved by showing that the degradation product of the generic is significantly more than 1% after being stored for 12 months at room temperature and shading.

The announcement also mentioned that the patentee Chugai is going to appeal the decision. However, even if the appeal was filed immediately in April 2022, and could overturn at the IP Tribunal of the Supreme Court (where the appeal should be filed), it is very unlikely that the appeal decision may be issued within the 9-month windows to suspend the DMA process of Haihe at the NMPA, notwithstanding that the complaint was
filed in November 2021 at the earliest (see above). In fact, there had been no news regarding the appeal as of June 2021.

It is worth noting that the Beijing IP Court's announcement specifically recited "allowing Chinese people to have good and cheap drugs". This may reflect the attitude of the Chinese government and the courts on the purpose of the Chinese drug patent linkage system, which does not seem to be too encouraging to original drug developers. On the other hand, as the generic may have tried to avoid the scope of ‘777 by proving to have more degradation product after 12-months of storage, it is curious to see how such could be qualified as "good" drugs.

**CNIPA “first” drug patent linkage decision**
The drug at issue is oxycodone hydrochloride sustained-release tablets to relieve severe pain (like those because of cancer) of Purdue Pharma under the brand OxyContin (basically, an opioid painkiller). The generic applicant is Yichang Renfu Pharmaceutical Industry Limited Liability Company (Renfu, 宜昌人福药业有限责任公司).

The patents at issue are 201210135209.X, 201510599477.0, and 201010151552.4.

201210135209.X and 201510599477.0 belong to the same family of WO/2008/023261A1 with priority from US60/840,244, and therefore also have the same filing date of 24 August 2007 and full term expiry of 24 August 2027.

201010151552.4 belongs to a different family of WO/2005/097801A1, with full term expiry on 30 March 2025.

As mentioned above, the CNIPA ruled that the generic drug from Renfu does not fall with the scope of the patents at issue.

Renfu also challenged that the recordal of 201010151552.4 (in the drug patent database at NMPA) is incorrect, but this challenge was not recognized by the CNIPA, while the reason is not specified in the CNIPA's announcement above.
The main claims of 201210135209.X, 201510599477.0, and 201010151552.4 could be summarized as below, in addition to the active ingredient oxycodone hydrochloride:

<table>
<thead>
<tr>
<th>CN patent</th>
<th>Requirements in main claim</th>
</tr>
</thead>
<tbody>
<tr>
<td>201210135209.X</td>
<td>at least one polyethylene oxide having, based on rheological measurements, an approximate molecular weight of at least 1,000,000, of at least 80wt% of the composition</td>
</tr>
<tr>
<td>201510599477.0</td>
<td>at least one polyethylene oxide having, based on rheological measurements, an approximate molecular weight of at least 800,000, of at least 80wt% of the composition</td>
</tr>
<tr>
<td>201010151552.4</td>
<td>Limits the amount of 14-hydroxycodeinone, an undesirable precursor of oxycodone, to be less than 15ppm in the composition</td>
</tr>
</tbody>
</table>

Again, it may not be that difficult for the generic of Renfu to get around the claims of the above 3 three patents. For example, this could be achieved by not using polyethylene oxide at all or using another component that does not literally or by equivalence fall within the scope, and showing that the undesirable precursor 14-hydroxycodeinone presents in an amount of more than 15ppm in the composition.

**Observations from CN drug patent linkage decisions as of June 2022**

It is interesting to see that the challenges from the generics in China thus far (the above challenge from Renfu, the challenge from Haihe in the BJ IP Court decision, and the first type 4.2 declaration from Hansoh) all refer to generics having seemingly properties inferior to the respective original drugs, while from the Beijing IP Court announcement that the purpose of the CN drug patent linkage system is to provide good and cheap drugs for people in China according to the BJ IP Court decision (My head is spinning to understand how drugs having more degradation product or undesirable precursor could be good drugs).

Another point to note is that the CNIPA's announcement above on 25 April 2022 also mentioned that 39 drug patent linkage complaints have been accepted out of 59 filed thus far, a significant increase from the previous CNIPA's announcement in October 2021 (12 accepted out of 23 filed). This shows that the CNIPA is indeed becoming the more favorable forum for drug patent linkage disputes.

The first drug patent linkage decision with the generic drug decided to fall within the scope of the patent(s) at issue still awaits to be seen.
Comparison between the US and CN drug patent linkage system

Thanks to the suggestion from Carol Nielsen of Nielsen IP, and review and comments by Corey Salsberg of Novartis, the appendix of this guide has a table showing the differences between the US and China drug patent linkage system.

Conclusion

While the “CN Orange Book” is based on the US Orange Book (as required under the US-CN Phase-I Trade Agreements 2020), the Chinese system has the following characteristics:

a) The patent information in the Chinese system is much more extensive than that of the US.

b) There is no drug data exclusivity information in the Chinese system.

c) Action could only be taken against type 4.2 declaration (the generic drug does not fall within the scope of the patent(s)) in China.

Regarding c), this may be the substantive difference of the CN system from the US system. In essence, in China, there is nothing the DMAH could do to delay the drug marketing approval process at the NMPA if the generic drug applicant simply alleges that the patent at issue should be declared invalid, and does not even challenge the validity of the patent afterwards.

There have been many changes on the “CN Orange Book” since its establishment on 29 June 2021. For example, in the very beginning, the platform did not have different tabs for entries of chemical drug, biological drug, and Chinese medicine. Now individual tabs are available, making searching and counting much easier. Finally, all five types (types 1-3, 4.1, and 4.2) declarations have been announced.

If you are an innovative drug owner, I am not surprised that you have already registered your drug patent information on the “CN Orange Book” to deter generic drug marketing approval application.

For others, including myself heavily involved in drug freedom-to-operate work, the “CN Orange Book” is a mountain full of treasures. Once the Chinese drug name is known (hint: have a look at the drug innovator’s Chinese website), the grant patent publication, claim(s)
and subject matter(s) relevant (with clear indication on what claim is covering which subject matter), full term expiry, and invalidation information (if available) are all available in one place.

After the first batch of drug patent linkage complaints have been accepted by the CNIPA and the Beijing IP Court at least in October 2021, while the two decisions thus far could be made within the required 9-months window, these two decisions only favored the generics. However, there may be legitimate reasons for these two decisions, as the patents relied on by the DMAHs in these decisions may not be too difficult to get around. By contrast, as shown in the first 4.2 declaration, the generic specifically avoided the relative strong patent directed to the chemical structure of the active ingredient of the drug.

Another point to note is that while the Chinese government has the intention that the CN drug patent linkage system is to provide Chinese people with good and cheap drugs, the Chinese generics that have challenged the DMAH do not appear to produce drugs as good as the corresponding original versions.

Author: Toby MAK, Tee & Howe Intellectual Property Attorneys ©2022
### Appendix - Differences between the US and China drug patent linkage systems

#### Chemical drug

<table>
<thead>
<tr>
<th>Aspect</th>
<th>US</th>
<th>CN</th>
</tr>
</thead>
<tbody>
<tr>
<td>New drug exclusivity</td>
<td>5 years</td>
<td>None</td>
</tr>
<tr>
<td>Types of declarations</td>
<td>4 types of declarations</td>
<td>4 types of declarations as the US, but type 4 declaration is further subdivided into the following types 4.1 and 4.2 declarations:</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• 4.1. the patent should be declared invalid</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• 4.2. the generic drug does not fall within the scope of the patent</td>
</tr>
<tr>
<td>Notification to DMAH</td>
<td>Generic drug applicant to notify the drug marketing approval holder (DMAH) of the type 4 declaration within 20 days after receiving FDA’s acknowledgment that the Abbreviated New Drug Application (ANDA) has been received and is ready for substantive review.</td>
<td>DMAH to find out announcement of notice of type 4 declaration from the CN drug patent linkage platform.</td>
</tr>
<tr>
<td>Filing of drug patent linkage complaint</td>
<td>DMAH to file suit at a court within 45 days after receipt of notice from the generic drug applicant.</td>
<td>DMAH to file complaint at the Beijing IP Court or the CNIPA with 45 days after announcement of notice of type 4 declaration, but ONLY for type 4.2 declaration. That is, there is no delay of the drug marketing approval process if a type 4.1 declaration is filed.</td>
</tr>
<tr>
<td>Stay of generic drug marketing approval process after filing the drug patent linkage complaint</td>
<td>After filing of the suit, the DMAH has an automatic 30-month stay of regulatory approval of the ANDA (which begins on “the later date of receipt” of the notice letter sent by the generic drug applicant) by the DMAH.</td>
<td>After acceptance of the complaint, and <em>if and only if</em> a decision was made within 9 months from the date of acceptance of the complaint that the generic drug falls within the scope of the patent, the NMPA would initiate the drug marketing approval process 20 days before the patent(s) expired. Otherwise, the marketing approval process would proceed as normal, i.e., as if drug patent linkage protection does not exist.</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
</tbody>
</table>
| Types of patents that could be registered | • The drug substance (active ingredient), **including different crystal forms**  
• The drug product (formulation and composition)  
• Medical use | • The drug substance (active ingredient), **excluding different crystal forms**  
[Note: However, composition patent including the different crystal forms could be registered]  
• The drug product (formulation and composition)  
• Medical use |
| Information contained in the drug patent database | Only US patent numbers and the full-term expiry date are listed. | Extensive information is listed in the drug patent database as below:  
1) The granted patent itself is available for downloading at a link on the same page.  
2) The claim number(s), and the relevant subject matter, are clearly identified.  
3) The full-term expiry of the patent(s) is clearly identified.  
4) Whether the patent has survived through invalidation is also specified in the note section. |
<p>| Incentive to generic drug applicant | 180 days marketing exclusivity to the first applicant to file the ANDA. | 12-months exclusivity period to the first successful challenger who files a type 4.1 declaration and subsequently invalidates the patent successfully. |</p>
<table>
<thead>
<tr>
<th>Aspect</th>
<th>US</th>
<th>CN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biologic exclusivity</td>
<td>12 years of total market protection, including 4 years before a biosimilar application may be submitted to the FDA, and another additional 8 additional the application may be approved.</td>
<td>None.</td>
</tr>
<tr>
<td>Stay of generic drug marketing approval process</td>
<td>None</td>
<td>None.</td>
</tr>
</tbody>
</table>
| Types of patents that could be registered  | Any patents identified by the reference product sponsor to a biosimilar applicant during the “patent dance”, in which the reference product sponsor and the biosimilar applicant exchange information of patents that may be infringed, and may include multiple waves of litigation. | • Sequence structure of the active ingredient  
• Medical use |
| Incentive to generic biologic applicant    | 12 to 42 months for “interchangeable biosimilars” (with more information and data available showing the impact of switching such that the intervention of the health care professional is not required, with the first approved by the FDA on 28 July 2021), depending on whether infringement suit is filed, and if filed, the duration of the infringement suit | None.                   |
The author Toby Mak has a somewhat unique background as a registered Chinese patent attorney. Before obtaining the Chinese qualification, he was trained under the UK patent system, and has passed some of the UK patent attorney’s examination papers. Toby is the current vice-chair of the Asian Practice Committee of the IPO, the current co-chair of the IP Practice in the Far East Committee of the AIPLA, and the past co-chair of the IP Practice in China Committee of the AIPLA. He is heavily involved in drug freedom-to-operate matters due to a client in Hong Kong, which makes him work from home in Hong Kong, while his firm and most of his colleagues are in Beijing.