FDA CIRCULAR
No. 2014-027


I. RATIONALE

On 13 October 2014, Administrative Order No. 2014-0034 was issued to (a) update and streamline regulatory approaches in licensing of drug establishments, (b) provide faster access of drug products to the public; and (c) promote transparency through the universal use of electronic transaction.

In line with the new rules and regulations on the licensing of establishments classified as drug manufacturers and its subclass (packer/repacker/trader), the Food and Drug Administration (FDA) hereby prescribes the requirements for application for initial and renewal issuance of License to Operate (LTO), variations, as well as other guidelines relevant to these establishments.

II. LICENSE TO OPERATE (LTO) APPLICATIONS

A. Documentary Requirements

1) Application Form
A completely filled-out and notarized application form signed by the pharmacist and owner/authorized representative must be submitted.

2) Proof of Business Name Registration
A valid proof of business name registration must be submitted:
(a) For single proprietorship - Certificate of Business Registration issued by the Department of Trade and Industry (DTI)
(b) For corporation, partnership and other juridical person - Certificate of Registration issued by the Securities and Exchange Commission (SEC) and Articles of Incorporation
(c) For cooperative - Certificate of Registration issued by the Cooperative Development Authority and the approved by-laws
(d) For government-owned or controlled corporation - the law highlighting the provision creating such establishment
The proof of business name registration must specify the exact and complete address, e.g., unit number, floor, building, lot, block, phase, street, barangay, city/municipality, province, where applicable.

3) Credentials of the Pharmacist and Other Qualified Personnel
   The credentials of the identified pharmacist-in-charge for a specific activity must be submitted, which include:
   (a) Valid PRC ID
   (b) Certificate of Attendance to appropriate FDA Licensing Seminar
   (c) Resignation letter of the pharmacist from previous employer (if previously employed).

   The other qualified personnel shall be listed, which include the (1) production manager/head, (2) quality assurance manager/head, (3) quality control manager/head, (4) authorized person for batch release, and (5) pharmacovigilance officer. The credentials will not be submitted during application but may be verified during inspection.

4) Risk Management Plan
   A general Risk Management Plan (RMP) for the establishment must be submitted. The RMP shall contain details on how to identify, characterize, prevent or minimize risk relating to the products they engage with. These shall include pharmacovigilance activities and interventions of the establishment to manage the risks.

5) Location Plan
   A sketch of the location of the establishment must be submitted which shall be used for inspection purposes. This sketch must indicate clear directions with identified landmarks to locate the establishment.

   In addition, the Global Positioning System (GPS) Coordinates in decimal degrees (DD) [Latitude and Longitude] must be indicated in the submission.

6) Site Master File
   The Site Master File (SMF) must be submitted, in accordance with the latest edition of the Pharmaceutical Inspection Cooperation Scheme (PIC/s) – Good Manufacturing Practice (GMP).

7) Proof of Payment
   Proof of payment (e.g., official receipt or authorized bank payment slip) must be included as proof of filing of application.

8) Self-Assessment Toolkit
   To guide and facilitate the submission, a Self-Assessment Toolkit (SATK) must be submitted, which will also serve as the worksheet during evaluation of FDA.
The list of documentary requirements for initial and renewal applications of LTO, reissuance of lost or destroyed LTO, as well as voluntary cancellation is attached as Annex A.

B. **Evaluation of Application**

1) Desktop Evaluation

All applications shall be initially reviewed by the respective FDA Regional Field Offices to determine compliance with the administrative and technical requirements.

The FDA, in the course of its evaluation may require additional or supplemental documents as proof of compliance to the existing regulations.

2) Pre-opening Inspection

After evaluation of the LTO application, the establishment shall be subjected to pre-opening inspection to determine compliance with the existing guidelines on Pharmaceutical Inspection Cooperation Scheme (PIC/S)-Good Manufacturing Practices (GMP).

In addition to the documentary requirements submitted during application (Section II, A of this Circular), the following documents shall be verified during inspection:

- Quality Management System
- Quality Manual and Standard Operating Procedures
- Contract Agreement (e.g., Manufacturing/ Packing / Repacking Agreement for Manufacturer-Trader, Packing for Manufacturer-Packer, Repacking for Manufacturer-Repacker)
- Qualification and Validation Documents
- Master and Batch Production Records
- Specifications
- Credentials of other qualified personnel
- Proof of Ownership/Lease Agreement of the space/bldg. by the establishment occupied
- Relevant reference materials (e.g., Republic Acts, PIC/s-GMP Guide, standard practice guidelines)
- Other procedures, protocols, records, and reports as required by PIC/s-GMP

The abovementioned additional documents will serve as proof of compliance by the establishment with the existing regulations on licensing.

A report shall be issued to the drug establishment after inspection, which shall be the basis for further decision/action of FDA (e.g.,
approval/disapproval of an application for LTO, and/or for such other purposes).

C. Post-licensing Inspection

All drug manufacturers and its subclass with approved LTO shall be subjected to routine inspection following the applicable provisions mentioned for pre-opening inspection under Section II, B of this Circular. In addition, major variation applications may require post-licensing inspection prior to the approval of such variation. Drug manufacturers and its subclass which are subject to regulatory action due to different triggers (e.g., violation of any of the provisions of FDA laws, rules and regulations, and any other laws related thereto, occurrence of adverse drug reactions, as well as other quality, safety, and/or efficacy issues) shall also be subject inspected.

D. Application for Variation

The following are the applicable variations to an approved LTO as drug manufacturer and its subclass:

1) Major Variation
   (a) Change of Ownership
   (b) Additional Production Line (e.g., sterile, penicillin)
   (c) Transfer of Location
   (d) Change of Activity

2) Minor Variation – Prior Approval
   (a) Expansion of Establishment
   (b) Change of Business Name
   (c) Zonal Change in Address

3) Minor Variation – Notification
   (a) Change of Pharmacist or other Qualified Personnel
   (b) Deletion of Activity
   (c) Transfer/Addition of Warehouse

FDA should be duly informed of any changes to the approved LTO, whether or not these are classified as variations described above. Other changes may also be added to the variations mentioned, which shall be subject of an appropriate regulation.

The list of documentary requirements for the abovementioned variations is attached as Annex B.

All variations are subject to the existing variation/amendment fee except for transfer of location which is subject to initial payment for two (2) years validity of LTO.

Drug manufacturer and its subclass applying for minor variations may continue business operations provided that an application for such variation has already been filed.
E. Accessibility

All electronic fillable forms shall be made accessible at the FDA Website.

III. EXEMPTION IN PROVIDING ANOTHER PHARMACIST FOR ADDITIONAL LICENSED ACTIVITY

Exemption may be granted to Drug Manufacturer-Repacker and Drug Manufacturer-Trader pharmacist to handle another non-manufacturing activity except retailing, provided that the following conditions are satisfied:

1) The activities sought to be licensed belong to one establishment only; establishment herein shall refer to a single business entity with the same business name registration and ownership who may engage in more than one licensed business activity;
2) The activities to be handled by the pharmacist are confined only in one office and warehouse within the same premises.

This exemption shall be duly noted in the application indicating the pharmacist’s duties and responsibilities as well as the schedule and hours of supervision to each establishment signed both by the pharmacist-in-charge and the owner/authorized representative.

IV. RESPONSIBILITIES OF OTHER IMPLEMENTING OFFICES

Consistent with the regulatory powers provided under (3), c, Sec. 2, Article III, Book I of the implementing rules and regulations of Republic Act No. 9711, FDA and its Regional Field Offices through the Director General may call on the assistance of any department office and/or government agency for the effective implementation of its rules and regulations.

In addition, Local Government Units (LGUs) are enjoined in monitoring licensed drug manufacturers in their localities for their compliance to the existing laws and their respective rules and regulations. Any violation found by the LGU inconsistent with the FDA rules and regulations shall be reported to FDA for regulatory action.

V. TRANSITORY PROVISIONS

Existing licensed establishments are required to submit their Risk Management Plan and GPS Coordinates upon renewal of their LTO.

VI. REPEALING CLAUSE/SEPARABILITY CLAUSE

Provisions of previous FDA circulars and memoranda that are inconsistent with this issuance are hereby withdrawn, repealed, and/or revoked accordingly. In case any part, term or provision of this FDA Circular is declared contrary to law or unconstitutional, other provisions which are not affected remain in force and effect.
VII. EFFECTIVITY

This Circular shall take effect upon approval and signature by the FDA Director General.

ATTY. NICOLAS B. LUTERO III, CESO III
Assistant Secretary of Health
OIC, Food and Drug Administration
ANNEX A

LIST OF DOCUMENTARY REQUIREMENTS FOR DRUG MANUFACTURER LTO APPLICATIONS

A. Initial LTO Application
   1) Application Form
   2) Proof of Business Name Registration
   3) Credentials of Pharmacist
   4) Risk Management Plan
   5) Location Plan
   6) Site Master File
   7) Proof of Payment (e.g. official receipt or authorized bank payment slip)
   8) Self-Assessment Toolkit

B. Renewal LTO Application
   1) Application Form
   2) Copy of Certifications issued as a result of LTO Variation
   3) Proof of Payment (e.g. official receipt or authorized bank payment slip)
   4) Self-Assessment Toolkit

C. Reissuance of Lost or Destroyed LTO
   1) Letter of Request
   2) Affidavit of Loss or Destruction
   3) Proof of Payment (e.g. official receipt or authorized bank payment slip)

D. Voluntary Cancellation of LTO
   1) Letter of Request
   2) Original LTO
ANNEX B

LIST OF REQUIREMENTS FOR VARIATION APPLICATIONS FOR DRUG MANUFACTURERS

A. Major Variations

<table>
<thead>
<tr>
<th>Change of Ownership</th>
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</thead>
<tbody>
<tr>
<td>C</td>
<td>There is a change of ownership of the drug establishment licensed.</td>
</tr>
<tr>
<td>D</td>
<td>1. Application Form</td>
</tr>
<tr>
<td></td>
<td>2. Proof of business name registration reflecting the name of new owner</td>
</tr>
<tr>
<td></td>
<td>3. Deed of sale or transfer of rights/ownership</td>
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<tr>
<td></td>
<td>4. Proof of payment</td>
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<td></td>
<td>5. Self-Assessment Toolkit</td>
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<table>
<thead>
<tr>
<th>Additional Production Line</th>
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<tbody>
<tr>
<td>C</td>
<td>An additional production line is an added type or class of products produced within the same manufacturing site (e.g., sterile line, penicillin line, etc.)</td>
</tr>
<tr>
<td>D</td>
<td>1. Application Form</td>
</tr>
<tr>
<td></td>
<td>2. Updated Site Master File</td>
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<tr>
<td></td>
<td>3. Proof of payment</td>
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<tr>
<td></td>
<td>4. Self-Assessment Toolkit</td>
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<table>
<thead>
<tr>
<th>Transfer of Location</th>
<th></th>
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<tbody>
<tr>
<td>C</td>
<td>1. Physical transfer of the drug establishment with changes in the previously approved address.</td>
</tr>
<tr>
<td></td>
<td>2. Other variations (e.g., change of pharmacist or key personnel, and/or business name) may also be included as long as the variation is noted in the application and the corresponding requirements for such changes are included. The payment remains as initial fee, regardless of the additional variation.</td>
</tr>
<tr>
<td>D</td>
<td>1. Application Form</td>
</tr>
<tr>
<td></td>
<td>2. Proof of business name registration reflecting the new address</td>
</tr>
<tr>
<td></td>
<td>3. New Location Plan</td>
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<tr>
<td></td>
<td>4. Updated Site Master File</td>
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<tr>
<td></td>
<td>5. Proof of payment</td>
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<td></td>
<td>6. Self-Assessment Toolkit</td>
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<tr>
<td>Change of Activity</td>
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<td>-------------------</td>
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<tr>
<td>C</td>
<td></td>
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<tr>
<td>1. Shall refer to an additional activity engaged by the manufacturer (e.g. LTO as Manufacturer with additional activity as Repacker)</td>
<td></td>
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<tr>
<td>2. Shall also refer to a change from the initially licensed activity (e.g. LTO as Manufacturer-Repacker to Manufacturer-Packer).</td>
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<tr>
<td>D</td>
<td></td>
</tr>
<tr>
<td>1. Application Form</td>
<td></td>
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<tr>
<td>2. Updated Site Master File</td>
<td></td>
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<tr>
<td>3. Proof of payment</td>
<td></td>
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<tr>
<td>4. Self-Assessment Toolkit</td>
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</tbody>
</table>
### B. Minor Variations – Prior Approval

#### Expansion of Establishment

| C | 1. Shall refer only to the expansion made which is adjacent to the existing location of the establishment and no additional product line involved.  
2. For Manufacturer-Trader, expansion shall include additional floors where the building is occupied. |
|---|---|
| D | 1. Application Form  
2. Updated Site Master File  
3. Proof of payment  
4. Self-Assessment Toolkit |

#### Change of Business Name

| C | 1. Change only in the business name  
2. No transfer of location or change of ownership. |
|---|---|
| D | 1. Application Form  
2. Proof of business name registration reflecting the new name of the drug establishment  
3. Proof of payment  
4. Self-Assessment Toolkit |

#### Zonal Change in Address

<table>
<thead>
<tr>
<th>C</th>
<th>Shall refer to change of the name/number of the street/building without physical transfer of the establishment.</th>
</tr>
</thead>
</table>
| D | 1. Application Form  
2. Document issued by the local municipality as proof of zonal change  
3. Proof of payment  
4. Self-Assessment Toolkit |
C. Minor Variations – Notification

<table>
<thead>
<tr>
<th>Change of Pharmacist or other Qualified Personnel</th>
</tr>
</thead>
</table>
| C | 1. There is a change of the identified pharmacist or other qualified personnel registered with FDA (e.g. production manager, quality assurance manager, quality control manager, authorized person for batch release)  
2. For other pharmacists and key personnel employed but not registered with FDA, changes on such shall not be required to apply for variation. |
| D | 1. Application Form  
2. Credentials (for change of pharmacist only)  
3. Proof of payment  
4. Self-Assessment Toolkit |

<table>
<thead>
<tr>
<th>Deletion of Activity</th>
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<tbody>
<tr>
<td>C</td>
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</tbody>
</table>
| D | 1. Application Form  
2. Termination of contract or conformance letter  
3. Proof of payment  
4. Self-Assessment Toolkit |

<table>
<thead>
<tr>
<th>Transfer/Addition of Warehouse</th>
</tr>
</thead>
</table>
| C | 1. Shall refer to the physical transfer of warehouse.  
2. Shall also refer to an addition of warehouse aside from the existing and previously inspected warehouse by FDA. |
| D | 1. Application Form  
2. New Location Plan  
3. Proof of payment  
4. Self-Assessment Toolkit |

*C – Condition  
*D – Documents required