

MOHAP e-Transformation

Site Registration - External User Guide

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1. Brief Overview

This manual is prepared to navigate applicants through the site registration process in Ministry of Health and Prevention (MOHAP) Portal.

This manual is designed to help applicant:

- Apply for Site Registration services
- Manage Site Registration applications

After completing this manual, the applicant should be able to perform all activities related to site registration on MOHAP Portal.

Please note: The Site Registration services are only available for the following Applicant Types:

Applicant Type	Rules
Agent	Should have un-expired license Can register any manufacturing site that is not local Can renew registration for any manufacturing site that is not local Can request minor variation for any manufacturing site's registration that is not local
Scientific Office	Should have un-expired license Can register any manufacturing site that is not local Can renew registration for any manufacturing site that is not local Can request minor variation for any manufacturing site's registration that is not local
Local Manufacturer	Should have un-expired license Can register any manufacturing site that is not local Can renew registration for any manufacturing site that is not local Can request minor variation for any manufacturing site's registration that is not local

Table 1: Applicant Types

2. Apply for Service

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Portal users who have logged in successfully to MOHAP portal will be directed to Dashboard screen where they can apply to MOHAP's e-services. This user manual will focus on the following e-services:

- 1. New Manufacturer Site Registration
- 2. Renewal of Registration to a Registered Site
- 3. Minor variation to Registered Site

For applicants to apply to their available services, they must go through the following process:

- 1. Create a new application
- 2. Complete application
- 3. Review application
- 4. Proceed to checkout

The sections below will help user through the step-by-step process of applying to an application.

Please note: Business Account users that are licensed by MOHAP will **NOT** be able to submit new applications if their license has **expired** but will be able to still view and apply actions for existing applications.

UNITED ARAB EMIRATES MINISTRY OF HEALTH & PREVENTION				العربية 💶	Registered Products
Dashboard All Applications			Open Registe	red Manufacture Sites	Manufacture Sites
	Dashboard			► New Application	Create a New Application
	36	170	85	2	
	Pending Correction	Pending with MOH	Pending Payment	Rejected	
	Applications Statistics		Renews	vals Expiry Date	
	Pending Correction	36 12%	Seleniu _15510	umNewPackName 050904330 25/02/2012	
			Mamdo	ouh_GSL_01 21/02/2014	
	 Pending with MOH 	170 58% 29	3 Pack N	Vame 23/04/2014	
	Pending Payment	85 29%	Cations Mamdo	ouh_Pack_03 15/05/2014	
	Rejected	2 0%	etisalat	t pack1 24/04/2017	
			Show n	more	

Figure A: Dashboard screen

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2.1 Create a New Application

The Applicant can request the registration, renewal or minor Variation of a registered site by either:

- 1. Selecting any of the following services from the Service Catalogue in MOHAP's website:
 - Registration of A Manufacturer of pharmaceutical Products or Medical devices
 - Renewal of Registration of A Manufacturer of pharmaceutical Products or Medical devices
 - Issuing The Certificate of Amendment of Any Registration Data of a Medical Company or a Factory That Has the Right to Marketing
- Creating a New Application form from the Applicant Portal and selecting any of the following services under the Site
 Registration services:
 - Site Registration
 - Site Minor Variation
 - Site Renewal

To create a new application form from the Applicant Portal, the user must do the following:

- a. Click on one of the **New Application** buttons found on the Dashboard Screen (or All Applications Screen) which opens New Application Screen.
- b. Pick one of the services found on New Application screen.
- c. Click Create Application button

New Ap	plication	×		
Drug	+			
Site	-			
Regi	istration —			
[Site Registration			
	Site Minor Variation			
	Site Renewal			
	Create Application	-	Create a New Application	
	Figure B: New Application Screen			

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Once the user selects a service and creates an application, the user will be redirected to Application Details screen.

2.2 Complete Application

To complete a site application, the applicant must fill the following three sections:

- 1. Site Details
- 2. Attachments

For an applicant to move from one section to another, they are required to fill the section's mandatory fields and tables.

2
Attachments

```
Figure C: Progress Bar
```

At any application stage, the applicant can perform the following actions:

- Save application
- Discard application
- Proceed to next section
- Go back to previous section

Warnings:

- In case the applicant did **not** complete a mandatory filed, a warning message marked in red will be displayed under the field text box highlighting the note: This Field is Required.
- In case the applicant did **not** fill a table, a warning message marked in red will be displayed under the table box highlighting the note: Table Cannot be empty.
- In case the applicant inputs a non-numeric character in a field that only accepts numbers, a warning message marked in red will be displayed under the field text box highlighting the note: This field accepts numbers only.

Display icons:

- 🔲 allows user to Delete table (usually found under Actions)
- Allows user to Edit table (usually found under Actions)

- 🔲 allows user to Add table
- 📙 allows the user to Save application
- 🖉 allows user to Insert attachment
- allows user to Insert photo

Tips: Each application section will contain a progress bar that shows the input progress of the application data

2.2.1 Site Registration

This service allows the applicant to register a new manufacturer site.

Once the applicant selects Site Registration as a service, they will be redirected to Site Registration – Site Details screen.

2.2.1.1 Site Details Screen

In Site Details screen, the applicant is required to fill the following sections:

- a. Site Details
- b. Product Classes
- c. Site Contact Person
- d. Market Authorization Holder
- e. Manufacturer Site License
- f. GMP Certifications*
- g. Activities Carried Out
- h. Manufacturing Lines
- i. Product Categories
- j. Quality Certifications
- k. Other Manufacturing Sites (Optional)
- I. Quality Management System (Optional)
- m. Countries Where Devices Are Approved and Sold (Optional)
- n. Documentation Procedures (Optional)

Rule: *Mandatory if the Drug Product Class is either: Conventional or Conventional Veterinary

Site Details	•	
--------------	---	--

Figure D: Dropdown Section List

Tip: Each section will contain a dropdown list of application section that will help users navigate between different sections

2.1.1.1.1 Site Details

In this section, the applicant is required to fill the following fields related to the site:

- Site Name
- Address
- P.O Box (Optional)
- City
- Country
- Phone
- Mobile (Optional)
- Email
- Fax (Optional)
- Website (Optional)
- Remarks (Optional)

	Site Details		T
	Site Details		
	1/14		
	Site Name ③		
	Site Name		
	Country	City	
	·	City	
	Address	P.O Box (Optional)	
	Address	P.O Box	
	Mobile (Optional)	Phone	
	Mobile	Phone	
	Email	Website (Optional)	
	Email	Website	
	Fax (Optional)		
	Fax		
	Remarks (Optional)		
	Remarks		
Discard			
Discald			

Figure E: Site Details Screen

2.1.1.1.2 Product Classes

For an applicant to add a new product class to their list of product classes, they must perform the following:

- a. Click on Add Product Class button
- b. Add the following fields:
 - Product Class
 - Sub-class (Optional)
- c. Click Add button

4% Completed				
	1 Site Details	2 Attachments		
	(i) All fields are mandatory, except for those labeled as optional.			
	Product Classes -			
	Product Class			
	Build your Product Class List	Add Product Class		
Discard		<	Back Next >	Click to Proceed

Figure F: Product Classes Screen

2.1.1.1.3 Site Contact Person

In this section, the applicant is required to fill the following fields related to the site contact person:

- Contact Name
- Address
- City
- Country
- Telephone
- Mobile (Optional)
- Fax (Optional)
- Email
- Website (Optional)

Site Details	Attact	hments
(i) All fields are mandatory, except	for those labeled as optional.	
Site Contact Person -		
3/14		
Name (i)	Address ①	
Name	Address	
City 🛈	Country ()	
City		~
Phone (i)	Mobile (Optional) ③	
Phone	Mobile	
Fax (Optional) (i)	Email 🚯	
Fax	Email	
Website (Optional) (i)		
Website		

Figure G: Site Contact Person Screen

2.1.1.1.4 Market Authorization Holder

In this section, the applicant is required to fill in the Market Authorization Holder details. To do that, the applicant must first search the MAH Name in the search bar and check if the MAH was previously registered.

If the MAH was not recognized, the applicant is asked to perform the following:

- 1. Fill the following MAH Details:
 - a. MAH Name
 - b. Address (Optional)
 - c. PO Box (Optional)
 - d. Country
 - e. City
 - f. Telephone

11 of

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- g. Email
- h. Fax (Optional)
- 2. Fill the following MAH Contact Person details:
 - a. Name
 - b. Address
 - c. City
 - d. Country
 - e. Telephone
 - f. Email
- 3. Fill the MAH's Financial Information (Optional)
 - a. Date of Establishment
 - b. Capital in US \$
 - c. Revenue in US \$
 - d. Year

If the MAH was recognized, the applicant must click on the MAH Name – which auto populates the MAH Details and the MAH Contact Person Section.

Once the applicant provides the section's required information, they can move to the next section by clicking **Next** button.

11% 0	Completed			
	3 Site Details	2 Attachments		
	All fields are mandatory, except for those labeled as optional.			
	Market Authorization Holder			
	4/14			
	Market Authorization Holder			
	Market Authorization Holder			
Discard		<	Back Next >	Click to Proceed
	Figure H: MAH Screen			

2.1.1.1.5 Manufacturer Site License

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In this section, the applicant is required to fill the following fields related to the Manufacturer Site License:

- Certificate Name
- Certificate Type
- Country/Authority
- Certificate Number
- Issue Date
- Expiry Date

Once the applicant provides the section's required information, they can move to the next section by clicking **Next** button.

14% Completed				
1 Site Details			2 Attachments	
(i) All fields are mandatory, except for	those labeled as o	ptional.		
Manufacturer Site Licens	ie -			
5/14				
Certificate Name (i)		Certificate Type (i)		
Certificate Name		Certificate Type		
Country/Authority (i)		Certificate Number (i)		
	~	Certificate Number		
Issue Date (i)		Expiry Date (i)		
dd/mm/yyyy	× 🗰	dd/mm/yyyy	× 🗐	

Figure I: Manufacturer Site License Screen

2.1.1.1.6 GMP Certifications*

In this section, the applicant is required to build their GMP (Good Manufacturing Practice) Certification details.

For an applicant to add a new GMP Certificate, they must perform the following:

a. Click Add Certificate button

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b. Fill the following fields:

- GMP Certificate Issuer
- Certificate Number
- Issue Date
- Expiry Date (Optional)
- GMP Certification Attachment (Optional)
- GMP Contact Person (Optional)
 - Contact Name
 - Address
 - City
 - Country
 - Telephone
 - Mobile
 - Fax
 - o Email
 - Website
- c. Click Add button

Rule: *This section is only mandatory for Drug Product Class: Conventional or Conventional Veterinary

Once the applicant provides the section's required information, they can move to the next section by clicking **Next** button.

	18% Completed			
	1 Site Details	2 Attachments		
	 All fields are mandatory, except for those labeled as optional. 			
	GMP Certifications -			
	6/14			
	Build your GMP certification Details List	I Add Certificate		
Discard		<	Back Next >	Click to
				Proceed

14 of

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2.1.1.1.7 Activities Carried Out

In this section, the applicant is required to select the activities carried out from the list provided.

Once the applicant provides the section's required information, they can move to the next section by clicking **Next** button.

21% Completed	
0	2
Site Details	Attachments
 All fields are mandatory, except for those labeled as optional. 	
Activities Carried Out -	
7///	
//14	
Activities (Optional)	
Manufacturing/Processing Of Starting Material	
Assembly	
Manufacture Of Dosage Forms	
Packaging & Labeling	
Storage & Handling	
Laboratory Testing	
N/A	
Batch Releaser (Certification)	

Figure K: Activities Carried Out Screen

2.1.1.1.8 Manufacturing Lines

In this section, the applicant is required to build their Manufacturing Lines list.

For an applicant to add a Manufacturing Line, they must perform the following:

- a. Click the Add Manufacturing Line button
- b. Fill the following fields:
 - Line Category
 - Line Subcategory
- c. Click the **Add** button

Once the applicant provides the section's required information, they can move to the next section by clicking **Next** button.

	25% Completed			
	1 Site Details	2 Attachments		
	All fields are mandatory, except for those labeled as optional.			
	Manufacturing Lines -			
	8/14			
	Manufacturing Lines			
	Build your Manufacturing Lines List	Add Manufacturing Line		
Discard		<	Back Next >	Click to Proceed

Figure L: Manufacturing lines Screen

2.1.1.1.9 Product Categories

In this section, the applicant is required to select the product categories from the list provided.

	29% Completed			
	3 Site Details	2 Attachments		
	(i) All fields are mandatory, except for those labeled as optional.			
	Product Categories -			
	9/14			
	Product Categories			
	N/A			
	Betalactam			
	Nonhazard			
	Cyctotoxic			
	Veterinary Use			
	Cytostics			
	Human Use			
	Hazard			
	Medicine .			
	Medical Device			
Discard		<	Back Next >	Click to

Figure M: Product Categories Screen

2.1.1.1.10 Quality Certifications

In this section, the applicant is required to build their Quality Certifications list.

For an applicant to add a Quality Certificate, they must perform the following:

- a. Click the Add Quality Certificate button
- b. Fill the following fields/attachments:
 - Certificate Issuer
 - Certificate Number
 - Issue Date
 - Expiry Date (Optional)
 - Attachment (Optional)
- c. Click the **Add** button

	32% Completed			
	1 Site Details	2 Attachments		
	All fields are mandatory, except for those labeled as optional.			
	Quality Certifications -			
	10/14			
	Quality Certifications			
	Build your Quality Certifications List	Add Quality Certificate		
Discard		<	Back Next >	Click to Proceed

Figure N: Quality Certifications Screen

2.1.1.1.11 Other Manufacturing Sites (Optional)

For an applicant to add a manufacturing site to the list, they must perform the following:

- a. Click the Add Other Site button
- b. Fill the following fields:
 - Company Name
 - City
 - Country
 - Operations Carried Out
- c. Click the Add button

	36% Completed		
	Site Details	2 Attachments	
	(i) All fields are mandatory, except for those labeled as optional.		
	Other Manufacturing Sites -		
	11/14		
	Other Manufacturing Sites		
	Build your Other Manufacturing Sites List	Add Other Site	
Discard		< Back Next >	Click to Proceed

Figure O: Other Manufacturing Sites Screen

2.1.1.1.12 Quality Management System (Optional)

For an applicant to complete the Quality Management System section, they must perform the following:

- a. Select the Type Of An Established Quality Management System from the following:
 - Partial Quality Management System
 - Full Quality Management System
- b. Fill the following fields:
 - Name of Facility
 - Address
 - City
 - Country
 - P.O Box

except for those labeled as a nt System – Management System (Optio System stem (Design, Production P iource Any Process (E.G., [Attach optional. nal) ostj Design & Development, Manufacturing, Warehou	ising,
except for those labeled as of nt System ~ Management System (Optio System stem (Design, Production P iource Any Process (E.G., D	optional. nal) ost) osesign & Development, Manufacturing, Warehou	ising,
nt System – Management System (Optio System stem (Design, Production P iource Any Process (E.G., D	nal) ost) Jesign & Development, Manufacturing, Warehou	ising,
Management System (Optio System stem (Design, Production P iource Any Process (E.G., E	nal) ost) design & Development, Manufacturing, Warehou	ising,
Aanagement System (Optio System Stem (Design, Production P iource Any Process (E.G., [nal) ost) Design & Development, Manufacturing, Warehou	ising,
	PO.BOX (Optional) (i)	
	PO.BOX	
	City (Optional) ③	
Ŧ	City	
	Ť	City (Optional) ③

Figure P: Quality Management System Screen

2.1.1.1.13 Countries Where Devices Are Approved and Sold (Optional)

For an applicant to add a country where devices are approved and sold, they must perform the following:

- a. Click the **Add Country** button
- b. Fill the following fields:
 - Country
 - Device Name
 - Authority that issues approval for marketing
- c. Click the **Add** button

0	2
Site Details	Attachments
() All fields are mandatory, except for those labeled as optional.	
Countries Where Devices Are Approved and Sold	•
13/14	
Country Where Devices Are Approved and Sold	
Build your Country Where Devices Are Approved and Sold List	Add Country

Figure Q: Countries Where Devices are Approved and Sold Screen

2.1.1.1.14 Documentation Procedures (Optional)

In this section, the applicant is required to select the product categories from the list provided.

Once the applicant provides the section's required information, they can move to the next section by clicking **Next** button.

		46% Completed			
	1 Site Details		Attachments		
	(i) All fields are mandatory, except for those labeled	as optional.			
C	Documentation Procedures -			•	
	4/14 ocumentation Procedures (Optional) Distribution Record Adverse Event Report Complaint Handling Alert And Modification Recall				
Discard			<	Back Next >	Click to Proceed
	Figure R: Documer	ntation Procedure Screen			

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2.2.1.2 Attachments

This section is where an applicant can upload Attachments before proceeding to submit the application. Once the applicant is done with this section, they can move to the Review section by clicking **Review** button.

The user can add more attachments to their Drug Registration application by performing the following:

- 1. Enter the Document Name
- 2. Click the Add Attachment icon 🖽
- 3. Upload Attachment

Tip: To view the list of attachments, please refer to Appendix I

		50% Completed		
	Site Details	2 Attachments		
	(i) All fields are mandatory, except for those labeled as op	ptional.		
F	Required Attachments			
A L F E T P o o n	A Legalized Letter Issued By The Company On Its Original Letterhead, Signed And Stamped By The Responsible Person In The Company, Authorizing A Person Or A Local Stabilishment To Submit The Registration Files On Its Behait, of The Drug Control Department. (Optional) Passe attach a copy of A legalized letter issued by the company on its rignal letterhead, signed and stamped by the responsible person in the company, authorizing a person or a local estabilishment to submit the egistration files on its behalt, to the Dug Control Department.	Legalized Valid Manufacturing License Issued By The Competent Authonity In Country Of Origin (Optional) Pease attach a copy of Legalized valid Manufacturing License issued by the competent authority in country of origin		
N G N P e n	Votarized Copies Of Registration Certificates Or Evidence Of MP Certification From Other Countries Where The Anundarturs Bie Begistered (Optional) Please attach a copy of Notarized copies of registration certificates or widence of GMC certification from other countries where the nanufacture site is registered New File	Legalized Current GMP Certificate Issued By The Competen Authority In Country Of Origin (Optional) Please attach a copy of Legalized current GMP certificate issued by th competent authority in country of origin	t.	
C F	Duality Certification (Optional) lease attach a copy of Quality Certification Www File	ISO 13485 Certification (Optional) Please attach a copy of ISO 13485 certification		
L S F a	ist Of The Products Manufactured And/Or Assembly By The Site (Optional) Please attach a copy of List of the products manufactured and/or assembly by the site	Site Master File (Optional) Please attach a copy of Site Master file Concer File		
	Document Name	B		
scard		< Ba	ck Review >	Click to Proceed
	Figure S: Attac	hment Screen		

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2.2.2 Site Renewal

This service allows the user to renew an expired or soon to be expired* manufactured site.

Zheijang Guobang Pharm	aceutical Co. Ltd.					Q
, , , ,						
Name	Country	City	Email	Expiry Date	Actoin	
Zhejiang Guobang Pharmaceutical Co. Ltd.	China	China-312 369 Shangyu, Zhejiang Province	someone@someon e.com	2008-01-01	<u>چې</u>	Renew Sit
Showing 1 to 1 of 1 entries		Firet	1 Dect			

Figure T: Registered Manufacture Sites - Renewal

Once the applicant selects Site Renewal as a service, they will be redirected to Site Renewal – Site Details screen where they are required to complete a renewal application.

In the Site Renewal - Site Details screen, the applicant is required to:

- a. Search the Site Name in the search bar
- b. Select Site to be renewed
- c. Click the **Renew** button
- d. Update fields/attachments in the Site Registration form
- e. Review application
- f. Proceed to checkout

	0% Completed				
		Select	2 Site Details	3 Attachments	
		i All fields are mandato	ry, except for those labeled as optional.		
		Manufacturing Site Name THOROUGHBRED REMEDIES	S MANUFACTURING LTD.		
		Country Ireland City			
		KILDARE Issue Date (Optional) 01/01/2003			
		Expiry Date (Optional) 01/01/2008			
	Discard				Renew Site
		Figu	ire U: Site Renewal – Site Details so	creen	
Display ico	on: ⁵ all	lows the user	to undo change		

Rule: *Only sites with 6 months or less left to expiration can be renewed

2.2.3 Site Minor Variation

This service allows the user to modify/update the allowed the site details.

stered Manufacture Site Amomed Pharma GmbH	'S					Q
Name	Country	City	Email	Expiry Date	Actoin	Click to
Amomed Pharma GmbH	Austria	Vienna	regulatory@amome d.com	2024-03-13	/	Modify Site
Showing 1 to 1 of 1 entries			< 1 > Last			-

Figure V: Registered Manufacture Sites - MV

Once the applicant selects **Site Minor Variation** as a service, they will be redirected to Site Minor Variation screen where they are required to:

- a. Search the Site Name in the search bar
- b. Select Site to be modified
- c. Click the Modify button

- d. Modify fields/attachments in the Site Registration form
- e. Review application
- f. Proceed to checkout

Rule ID	Rules							
R1	For a site's details to be modified, the site must not be expired.							
R2	Users will be allowed to modify only certain open fields - not all fields are necessarily open for Variation							
	Table 2: Site Minor Variation Rules							
	0% Completed							
	Select Site Details Attachments							
	All fields are mandatory, except for those labeled as optional.							
	Manufacturing Site Name Marn_UAT_Expired Country							
	United Arab Emirates City Dubai							
	Issue Date (Optional) 29/03/2019 Expiry Date (Optional) 16/04/2049							
	Discard Click to Modify Site							
	Figure W: Site Minor Variation screen							
	Display icon: ⁵ allows the user to undo change							

2.3 **Review Application**

After the applicant completes filling all the Site Registration form, they must click on the **Review** button to proceed with application submission. Once the applicant clicks the **Review** button, they will be redirected to Review screen where they are able to perform the following actions:

- Edit Application
- ✤ Save Application

- Discard Application
- Go Back to Previous Section
- Read & Accept General Terms & Conditions
- Submit Application

< Back Review	
lease take a moment to check that everything is correct. You can edit anything that's not right	
> Product Details	
> New Pack Details	
> Attachments	
I Accept The General Terms And Conditions	
Discard	Subn

Figure X: Review Screen

Please note: to proceed with application submission, the applicant must read & accept the General Terms and Conditions.

Once the applicant reviews application and accepts the General Terms and Conditions, they must click the **Submit** button to proceed to checkout.

Warning: In case the user did **not** fill all mandatory fields, a warning message will be displayed noting that applicant must fill all required fields before submitting the application.

Display icon:

- allows user to Edit application details
- allows user to view table

2.4 Checkout

After an application has been submitted, the applicant will be redirected to Checkout screen where they will be asked to pay the following fees which will be based on the service the applicant applied for:

- > Application Fees*
- Processing Fee

To proceed with payment, the applicant must click on the **Checkout** button shown at the bottom-right corner of the Checkout screen which will redirect the applicant to MOHAP's Payment Gateway (e-dirham).

P	Irice	
	Request For Issuance / Renewal Of GSL Product 100	
	Registration Of New GSL Product 5000	
	Total 5100	
		Checkout

Figure Y: Checkout Screen

Rule: *Site Minor Variation service does not have an application fee.

Tip: Overrun Pop-up Blocker to open MOHAP's Payment Gaterway (e-dirham)

2.4.1 Site Registration

The Site Registration fees include the following:

- > Application Fee: AED 100
- Processing Feel AED 10,000

Once the payment is done successfully, the application will be sent to a MOHAP officer for review where they could perform the following:

- Issue Certificate
- Reject Application
- Send Back Application

2.4.2 Site Renewal

The Site Renewal fees include the following:

> Application Fee: AED 100

Processing Feel AED 10,000

Once the payment is done successfully, the application will be sent to a MOHAP officer for review where they could perform the following:

- ✤ Issue Certificate
- Reject Application
- Send Back Application

2.4.3 Site Minor Variation

The Site Registration fees only includes the Processing Fee: AED 2,000

Once the payment is done successfully, the application will be sent to a MOHAP officer for review where they could perform the following:

- Issue Certificate
- Reject Application
- Send Back Application

3. Take Required Actions

Once an application has been reviewed by a MOHAP officer, the officer may request the applicant to take actions before proceeding with application. The applicant will get notified via SMS or email once the officer requires an action.

For a user to view and take these actions, they must:

- a. Click on the **Application Number** in All Applications screen which redirects user to Submitted Application screen
- b. Click on Required Actions tab
- c. Click on the Action which opens an Action window

Take	Pending Correction	Application Type:Site Registration Submitted On:04/04/2019	
Required Actions	Application details Required actions 1 Certificates	Application History	
	New Site Details Attachments		

Figure Z: Submitted Application Screen

3.1 Application Correction

In case the MOHAP Officer identifies fields/attachments in application that require Variation, the officer will send the application back to applicant for correction.

For the applicant to view and change the fields/attachments that require correction, the applicant must:

- a. Click on **Correction** button found in Required Actions tab which redirects applicant to Correction screen
- b. Make changes to fields/attachments*
- c. Add comment in Comment box
- d. Click on Submit Correction button
- e. Confirm correction

Once the applicant confirms correction, the application will be sent back to officer for further review.

Rule ID	Rules
*R1	*Only fields/attachments needed for correction are displayed for change
R2	The application will be considered as rejected if the applicant did not send back corrected application within 60 days

Table 3: Correction Rules

Pending Correction	Application Type Site Registration Submitted On 04/04/2019		
Application details Required actions () Certificates	Application History		
SREG-2019-000207 Correction Apr 4, 2019			
Figure AA: Submitted Applicat 4/5 Unresolved Request(s)	tion screen – Pending Correction		
Figure BB: Field	Correction Cursor		
MAH Name 🛈	MAH Name 🛈		
Market Authorization Holder Nam	SAG Manufacturing S.L.U		
Figure CC: Modified Field	Figure DD: Unmodified Field		
Legalized Current GMP Certificate Issued By The Competent Authority In Country Of Origin (Optional) No file attached Legalized Valid Manufacturing License Issued By The Competent Authority In Country Of Origin (Optional) Attachment	Notarized Copies Of Registration Certificates Or Evidence Of GMP Certification From Other Countries Where The Manufacture. Site Is Renistered (Dotional). No file attached 11/1 Unresolved Request(s) A Legalized Letter Issued By The Company On Its Original Letterhead, Sign Person Or A Local Establishment To Submit The Registration Files On Its Behalf, To The Drug Control Department. (Optional) & Attachment	~	
~ Comments			
Comment			
Corrected!			
	Submit Corrections	 ↓	Click to Submit



Display icons: ^O allows the user to undo change

Tips:

- i. Field Correction Tool helps user navigate from one unsolved request to another
- ii. Fields required for change are highlighted in Red

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iii. Modified fields are highlighted in Yellow.

4. Rejected Applications

Once an application has been rejected by a MOHAP Officer, an email or an SMS will be sent to the applicant containing the application's latest updates and results.

To view the rejection comments given by MOHAP Officer and the step at which the application got rejected, please click on the **Application History** button.

			×	
	Application History	/		
	Date	Action	Comment	
	11-04-2019	Payment Received	Transaction was processed successfully.	
	11-04-2019	submit		
	11-04-2019	submit		
	11-04-2019	Pending/Delayed	Pending delayed 1 time	
	15-04-2019	Reject	Rejected by Officer Application does not meet MOHAP standards	
	Close			
	Figu	ire FF: Applicatio	on History	
ected				Application Type:Site Registra Submitted On:11/04/2
etails Required actions	0 Certificates			49 Application F
ite Details				

Figure GG: Submitted Application Screen – Application Details

5. Printouts

Attachments

Once an application has been reviewed and approved by a MOHAP officer, the officer will issue the applicant a Certificate associated with the service type. The applicant will get notified via SMS or email once the officer issues the certificate.

For a user to view and printout a certificate, they must:

a. Click on the **Application Number** in All Applications screen – which redirects user to Submitted Application screen

- b. Click on the Certificate tab
- c. Click on the Attachment which opens the certificate as a PDF
- d. Print PDF

	Issued			Application Type:Site Registration Submitted On:14/04/2019
Open Certificates	Application details Required actions Certificates		4 Application History	
	Certificate	Issue Date	Expiry Date	Print
	Site Registration	14/04/2019	13/04/2024	Attachment

Figure HH: Certificate Screen

The Site Registration Certificate will be valid for 5 years.

The Site Registration Certificate will contain the following information:

- Certificate Number
- > Registration Number
- Committee Meeting No
- Payment Receipt No.
- Manufacturing Site Name
- > Address
- ➢ First Reg. Date
- Reg. Expiry Date
- > Meeting Date
- > Payment Date
- > Activities Registered For
- > Non-Hazard Line(s) of Production Registered For
- > Manufacturing Site for product Class(s)

Rule: Site Variation will not affect the Site Registration's expiry date.

6. Appendix

Attachments that the applicant will upload when submitting a site registration application.

Documents	Mandatory
A legalized letter issued by the company on its original letterhead, signed and stamped by the responsible person in the company, authorizing a person or a local establishment to submit the registration files on its behalf, to the Drug Control Department.	Y
Legalized valid Manufacturing License issued by the competent authority in country of origin	Y
Legalized current GMP certificate issued by the competent authority in country of origin	N
Quality Certification	N
ISO 13485 certification	N
List of the products manufactured and/or assembly by the site	Ν
Site Master file	Ν
Notarized copies of registration certificates or evidence of GMP certification from other countries where the manufacture site is registered	N

Table 4: List of Attachments