



Asian Harmonization Working Party
WORKING TOWARDS MEDICAL DEVICE HARMONIZATION IN ASIA

FINAL DOCUMENT

Title: Post Market Resource Center

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1. Objectives

The Post Market Resource Center is a tool developed by Work Group 4 of AHWP to provide a "one-stop" location for Regulatory Authorities (RAs) and the Medical Device Industry (Industry) to access to post-market regulations and reporting information easily across the world.

1. Adverse Event Reporting

1.1 Reporting System

- (a) AHWP Members
 - (i) China (required access)
<http://114.255.93.220/sso/login?service=http%3A%2F%2F114.255.93.220%2FFP%2FcasAuthUser>
 - (ii) Chinese Taipei (required access)
<https://gms.fda.gov.tw/tcbw/>
http://www.fda.gov.tw/TC/siteContent.aspx?sid=4243#.V1TYX_I97Dc
 - (iii) Hong Kong SAR
<http://www.mdco.gov.hk/english/report/report.html>
 - (iv) Kingdom of Saudi Arabia
<http://ncmdr.sfda.gov.sa/>
 - (v) Malaysia
http://www.mdb.gov.my/mdb/index.php?option=com_content&task=view&id=13&Itemid=36
 - (vi) Republic of Korea
(required access)
<https://emed.mfds.go.kr/>
 - (vii) Singapore:
http://www.hsa.gov.sg/content/hsa/en/Health_Products_Regulation/Safety_Information_and_Product_Recalls/Report_Adverse_Events_related_to_health_products.html
- (b) GHTF Countries
 - (i) Australia
<https://www.tga.gov.au/medical-devices-safety>
 - (ii) Canada
<http://www.hc-sc.gc.ca/dhp-mps/medeff/report-declaration/index-eng.php>
 - (iii) EU
 - A. France (French only)
<http://ansm.sante.fr/Declarer-un-effet-indesirable/Votre-declaration-concerne-un-dispositif-medical/Votre-declaration-concerne-un-dispositif-medical/Votre-declaration-concerne-un-dispositif-medical-Vous-etes-un-fabricant-un-distributeur>
 - B. Germany
http://www.bfarm.de/EN/MedicalDevices/vigilance/_node.html;jsessionid=72F58EBAF78E6D914DE67BA5A2BA5311.1_cid340
 - C. Switzerland
<https://www.swissmedic.ch/medizinprodukte/02675/02677/index.html?lang=en>
 - D. UK
<https://www.gov.uk/guidance/send-and-receive-information-on-adverse-drug-reactions-adrs>
 - (iv) Japan
<http://www.e-gov.go.jp/shinsei/index.html>
 - (v) United States
<http://www.fda.gov/Safety/MedWatch/default.htm>

1.2 Reporting Form

(a) AHWP Members

(i) China

<http://114.255.93.201/xzzx/>

(ii) Chinese Taipei

<http://www.fda.gov.tw/tc/includes/SiteListGetFile.ashx?mid=133&id=10510&chk=bae3b545-aea6-4307-928d-83ae4aab790b>

(iii) Hong Kong SAR

http://www.mdco.gov.hk/english/mdacs/mdacs_af/files/LRP_Adverse_Incident_Reporting_form.doc

(iv) Kingdom of Saudi Arabia

<http://ncmdr.sfda.gov.sa/>

(v) Malaysia

http://www.mdb.gov.my/mdb/index.php?option=com_content&task=view&id=19&Itemid=115

(vi) Republic of Korea



[서식_1]_의료기기
_이상사례_보고서(

(vii) Singapore

http://www.hsa.gov.sg/content/dam/HSA/HPRG/Medical_Devices/Updates_and_Safety_reporting/Adverse_Event_Reporting/Interactive_Industry%20Adverse%20Event%20Report%20Form_Aug%202015.pdf

(b) GHTF Countries

(i) Australia

<https://www.tga.gov.au/sites/default/files/devices-argmd-p3.pdf>

(ii) Canada

<http://www.hc-sc.gc.ca/dhp-mps/medeff/report-declaration/index-eng.php>

(iii) EU

<http://ec.europa.eu/DocsRoom/documents/15506/attachments/3/translations/en/renditions/native>

A. France (French only)

<http://ansm.sante.fr/Declarer-un-effet-indesirable/Votre-declaration-concerne-un-dispositif-medical/Votre-declaration-concerne-un-dispositif-medical/Votre-declaration-concerne-un-dispositif-medical-Vous-etes-un-fabricant-un-distributeur>

B. Germany

http://www.bfarm.de/EN/Service/Formulare/medDev/mp-forms-startseite_en.html

C. Switzerland

<https://www.swissmedic.ch/medizinprodukte/02675/02677/index.html?lang=en>

(iv) Japan

<http://www.pmda.go.jp/safety/reports/mah/0014.html>

(c) GHTF

<http://www.imdrf.org/documents/doc-ghtf-sg2.asp>

1.3 Guidance Notes

(a) AHWP

(i) China

<http://114.255.93.201/zcfg/ylqx/>

(ii) Chinese Taipei

<http://www.fda.gov.tw/TC/includes/GetFile.ashx?MID=133&id=28272&chk=0a6a912d-ee52-4e08-ad74-9d882c91aaba>

(iii) Hong Kong SAR ([GN-03] Guidance Notes for Adverse Incident Reporting by Local Responsible Persons)

http://www.mdco.gov.hk/english/mdacs/mdacs_gn/files/gn_03.pdf

(iv) Kingdom of Saudi Arabia

<http://ncmdr.sfda.gov.sa/>

(v) Republic of Korea

- **Reporting Manual for Health Care Professional, Industry, and Patient**
<http://www.mfds.go.kr/index.do?searchkey=title:contents&mid=695&searchword=부작용&cd=&pageNo=1&seq=11503&cmd=v>
- **Guidelines per specific product categories**
 - Soft Contact Lens, Filler, Cardiac Stent
<http://www.mfds.go.kr/index.do?searchkey=title:contents&mid=1161&searchDivision=의료기기&searchClass=&searchword=유해사례&searchSubDivision=&pageNo=1&seq=7285&cmd=v>
 - Breast Implant, Hip Implant, Knee Implant
<http://www.mfds.go.kr/index.do?searchkey=title:contents&mid=1161&searchDivision=의료기기&searchClass=&searchword=유해사례&searchSubDivision=&pageNo=1&seq=5997&cmd=v>

(vi) Singapore (GN-05: Guidance on the Reporting of Adverse Events for Medical Devices)

http://www.hsa.gov.sg/content/dam/HSA/HPRG/Medical_Devices/Overview_Framework_Policies/Guidances_for_Medical_Device_Registration/GN-05-R2_Guidance%20on%20the%20Reporting%20of%20Adverse%20Events%20for%20Medical%20Devices.pdf

(b) GHTF Countries

(i) Australia

<https://www.tga.gov.au/database-adverse-event-notifications-daen>

(ii) Canada

<http://www.hc-sc.gc.ca/dhp-mps/medeff/report-declaration/index-eng.php>

(iii) EU

https://ec.europa.eu/growth/sectors/medical-devices/guidance_en

(iv) Japan

<http://www.pmda.go.jp/safety/reports/mah/0014.html>

(v) United States

<http://www.fda.gov/Safety/MedWatch/default.htm>

2. Safety Information

2.1 Field Safety Information Reporting

(a) AHWP Members

- (i) China (each CFDA at province level post its own information, a few key provincial CFDA website listed as follows:)

<http://www.sda.gov.cn/WS01/CL0861/>

<http://www.shfda.gov.cn/gb/node2/yjj/aqgz/cpzh/ylqxcpszdh/n5110/index.html>

- (ii) Chinese Taipei

<http://www.fda.gov.tw/TC/siteList.aspx?sid=4275>

<http://www.fda.gov.tw/TC/site.aspx?sid=4232>

- (iii) Hong Kong SAR

<http://www.mdco.gov.hk/english/safety/safety.html>

- (iv) Kingdom of Saudi Arabia

<http://ncmdr.sfda.gov.sa/>

- (v) Malaysia



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- (vi) Republic of Korea

(required access)

<https://emed.mfds.go.kr/>

- (vii) Singapore

http://www.hsa.gov.sg/content/hsa/en/Health_Products_Regulation/Medical_Devices/Safety_reporting/Field_Safety_Corrective_Action.html

(b) GHTF Countries

- (i) Australia

<https://www.tga.gov.au/medical-devices-safety>

- (ii) Canada

<http://www.hc-sc.gc.ca/dhp-mps/medeff/report-declaration/index-eng.php>

- (iii) EU (Guidance Section 2.12 Market Surveillance)

(Guidance) https://ec.europa.eu/growth/sectors/medical-devices/guidance_en

(Form)

<http://ec.europa.eu/DocsRoom/documents/15506/attachments/5/translations>

A. France (French only)

<http://ansm.sante.fr/Declarer-un-effet-indesirable/Votre-declaration-concerne-un-dispositif-medical/Votre-declaration-concerne-un-dispositif-medical/Votre-declaration-concerne-un-dispositif-medical-Vous-etes-un-fabricant-un-distributeur>

B. Germany

(Guidance)

http://www.bfarm.de/EN/MedicalDevices/vigilance/_node.html;jsessionid=72F58EBAF78E6D914DE67BA5A2BA5311.1_cid340

(Form) http://www.bfarm.de/EN/Service/Formulare/medDev/mp-forms-startseite_en.html

C. Switzerland

<https://www.swissmedic.ch/medizinprodukte/02675/02677/index.html?lang=en>

D. UK

<https://www.gov.uk/government/publications/report-a-non-compliant-medical-device->

[enforcement-process](#)

- (iv) Japan
<http://www.pmda.go.jp/safety/reports/mah/0014.html>
- (v) United States
<http://www.fda.gov/Safety/Recalls/IndustryGuidance/ucm129259.htm>

2.2 Safety Alert Information

(a) AHWP Members

- (i) China
http://114.255.93.201/xxtb_255/ylqxbjsjxxtb/
- (ii) Chinese Taipei
<http://www.fda.gov.tw/TC/siteList.aspx?sid=4275>
<http://www.fda.gov.tw/TC/site.aspx?sid=4232>
- (iii) Hong Kong SAR
<http://www.mdco.gov.hk/english/safety/safety.html>
- (iv) Kingdom of Saudi Arabia
<http://ncmdr.sfda.gov.sa/>
- (v) Malaysia



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- (vi) Republic of Korea
<http://www.mfds.go.kr/index.do?mid=734>
- (vii) Singapore
http://www.hsa.gov.sg/content/hsa/en/Health_Products_Regulation/Medical_Devices/Safety_reporting/Field_Safety_Corrective_Action.html

(b) GHTF Countries

- (i) Australia
<https://www.tga.gov.au/alerts>
- (ii) Canada
<http://www.hc-sc.gc.ca/dhp-mps/medeff/report-declaration/index-eng.php>
- (iii) EU
 - A. France
(Safety Alert) <http://www.ansm.sante.fr/S-informer/Informations-de-securite-Autres-mesures-de-securite#dm>
(Recall) <http://www.ansm.sante.fr/S-informer/Informations-de-securite-Retraits-de-lots-et-de-produits#dm>
 - B. Germany
http://www.bfarm.de/SiteGlobals/Forms/Suche/EN/kundeninfo_Filtersuche_Formular_en.html?nn=3497208&searchEngineQueryString=search+item
 - C. Switzerland
https://www.swissmedic.ch/rueckrufe_medizinprodukte/index.html?lang=en
 - D. UK
https://www.gov.uk/drug-device-alerts?keywords=&alert_type%5B%5D=devices&issued_date%5Bfrom%5D=&issued_date%

[5Bto%5D](#)

(iv) Japan

<http://www.pmda.go.jp/english/safety/info-services/0014.html>

(Urgent Notice)

<http://www.pmda.go.jp/safety/info-services/devices/0092.html>

(Safety Information- Japanese)

<http://www.pmda.go.jp/safety/info-services/drugs/calling-attention/safety-info/0043.html>

(Safety Information – English)

<http://www.pmda.go.jp/english/safety/info-services/drugs/medical-safety-information/0002.html>

(v) United States

<http://www.fda.gov/safety/recalls/enforcementreports/>