## OFFICE OF PHARMACY AFFAIRS (OPA) 340B REGISTRATION FORM FOR OUTPATIENT FACILITIES USING MEDICARE COST REPORT

A complete registration package must include the information noted in sections I-VI below. In addition, the hospital may be required to provide additional supporting documentation including:

- 1. A copy of the latest filed Medicare cost report Worksheet S that includes a digital encrypted signature stamp\*;
- 2. A copy of Worksheet A, Worksheet C, Worksheet S-2 and Worksheet E Part A from the latest filed Medicare cost report (for the DSH adjustment percentage in II, A, below).
- 3. The associated trial balance.
- \* The date and time prepared listed in the upper right corner of all worksheets must match the date and time of the digital encrypted signature stamp.

The entire registration package must be submitted on the same day to be considered complete. A registration that is submitted without any of the required documentation will be rejected.

I. HospitalInformation:
Hospital (Main Provider) Name:
Hospital (Main Provider) Medicare Provider Number:
Hospital (Main Provider) Employer Identification Number:
Hospital (Main Provider) Street Address:
II. Hospital Outpatient Facility Information:
Please complete the following section and include all requested information. This registration form and supporting documentation must be completed and submitted according to the established deadlines that are published on the OPA website (www.hrsa.gov/opa). The registration process is not complete unless the registration form has been completed its entirety (all requested information is filled in on the form) and all required supporting documentation is submitted on the same day to OPA. <b>Incomplete packages will not be processed.</b>
Indicate the following regarding the outpatient facility to be registered:
Name of Facility:
Outpatient Employer Identification Number (if different from parent):
Facility Medicare Provider Number (if different):
Facility Street Address (P.O Boxes are not allowed):
Facility Billing Address (if different):
Facility Shipping Address (PO Boxes are not allowed):
City: State:ZIP:

## **III. Cost Center Information:**

Hospitals registering outpatient facilities must identify one or more specific cost center lines that the facility being registered falls under on the organization's most recently filed Medicare cost report. In the following fields, enter the Net Expenses for Allocation for the entire line (Worksheet A, Column 7) followed by the total outpatient charges for the entire line (Worksheet C, Column 7).

Next, enter expenses associated with the specific clinic, service or facility being registered – if more than one clinic, service or facility is rolled up to a single cost center (e.g., Line 90/Clinic), these figures will come from the corresponding working trial balance. (For cost centers/lines that reflect only a single outpatient clinic, service or facility, these figures will come directly from Worksheet A, Column 7 and will be the same as those provided for the entire line -- e.g., Line 90.xx subscripts). Finally, enter the outpatient revenue associated with the specific clinic or service being registered (also from the trial balance).

Line Number/Description	Net Expenses (Worksheet A)	Outpatient Charges (Worksheet C)	Specific Service/Clinic Cost (Trial Balance)	Specific Service/Clinic Outpatient Revenue (Trial Balance)

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At this site, v	will the covered	entity bill	Medicaid	fee-for-serv	rice for	drugs p	ourchased	at 340B	prices?
Yes □	No □	-							

If the answer is yes, please provide the state(s) and associated billing number(s) listed on the claims to bill Medicaid feefor-service for particular states that you plan to bill for 340B drugs in the space(s) below (this could include numbers for the state your hospital is located in and any out-of-state Medicaid agencies your hospital plans to bill for 340B drugs). All numbers you plan to use to bill Medicaid fee-for-service should be provided and may include the billing provider's national provider identifier (NPI) only, state assigned Medicaid number only, or both the NPI and state assigned Medicaid number. Do not list a state for which the covered entity will not bill Medicaid fee-for-service for drugs purchased at 340B prices.

HRSA exports the Medicaid billing information listed in this site's 340B OPAIS record to generate the quarterly Medicaid exclusion file (MEF). HRSA requires the information on the MEF be accurate and complete for every registered site in the 340B OPAIS, and that covered entities follow any additional state Medicaid requirements in order to prevent duplicate discounts.

While this site may request a change to its 340B OPAIS record at any time, the Medicaid fee-for service billing practice at this site, must match the quarterly MEF.

State	State Assigned Medicaid Number	NPI

All covered entities should notify OPA prior to any change in Medicaid billing status. For more information, please visit the HRSA website.

## V. Statutory Prohibition on Group Purchasing Organization Participation:

Section 340B(a)(4)(L)(iii) of the Public Health Service Act, which is reiterated in the Statutory Prohibition on Group Purchasing Organization Participation Policy Release (2013-1), requires that the hospital not obtain covered outpatient drugs through a group purchasing organization or other group purchasing arrangement. **This is a requirement for Disproportionate Share Hospitals, Children's Hospitals, and Free Standing Cancer Hospitals**.

The authorizing official must certify that this hospital will not participate in a group purchasing organization or group purchasing arrangement for covered outpatient drugs as of the date of this listing on the 340B OPAIS. If drugs are purchased using a GPO for covered outpatient drugs while participating in the 340B Program, the covered entity understands that this violates program eligibility requirements and that the covered entity is obligated to inform OPA and may be required to repay manufacturers for the 340B discount received.

		☐ Yes, I confirm
		□ Not applicable
VI. Signed Agreement:		
I acknowledge that I am familiar with the most rece Medicare certification of hospital components as of Pursuant to those guidelines, I request that the att OPAIS of 340B covered entities. I have examined covered entity's most recently filed Medicare cost Medicare provider number listed above. I further a published guidelines and regulations.	one cost center and HRSA's ached list of qualifying out the list and certify that eac report and is an integral pa	s guidelines for hospital outpatient facilities. patient facilities be added to the 340B th outpatient facility is reimbursable on the art of the aforementioned hospital under the
The undersigned represents and confirms that he/ and certifies that the contents of any statement ma undersigned further acknowledges the 340B cover	ade or reflected in this docu	ument are truthful and accurate. The
As an Authorized Official, I certify on behalf of the	covered entity and its outp	atient facilities that:
<ul><li>(1) all information listed on the 340B Program 340 correct;</li></ul>	B OPAIS for the covered e	entity will be complete, accurate, and
<ul> <li>(2) the covered entity will meet all 340B Program of Health Service Act when applicable, regarding covered entity hospital does not obtain covere group purchasing arrangement;</li> <li>(3) the covered entity will comply with all requirem accompanying regulations including, but not lind diversion (section 340B(a)(5)(A) and (B) of the</li> </ul>	the group purchasing orga d outpatient drugs through ents of Section 340B of the mited to, the prohibition ag	anization prohibition - which states that the a group purchasing organization or other e Public Health Service Act and any ainst duplicate discounts/rebates and
<ul> <li>(4) the covered entity will maintain auditable recomparagraph (3) above, pursuant to section 340E</li> <li>(5) the covered entity acknowledges its responsibile eligibility and/or breach by the covered entity of</li> <li>(6) the covered entity acknowledges that if there is covered entity may be liable to the manufacture and, depending upon the circumstances, may</li> <li>In addition, I have read all applicable registration reviewed if the required supporting documents</li> </ul>	B(a)(5)(C) of the Public Heality to contact OPA as soon of any of the foregoing; and a breach of the requirement of the covered outpatier be subject to removal from on instructions and I am as	alth Service Act; In as possible if there is any change in 340B Is ents described in paragraph (3) that the int drug that is the subject of the violation, in the list of eligible 340B entities.
Please provide any additional information that	may be helpful in reviewir	ng this registration for 340B eligibility:
Signature of Authorizing Official		Date
Name & Title of Authorizing Official and Title (please print or type)(e.g.CEO,CFO,COO)	Phone	Email

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information unless it displays a currently valid OMB control number. The OMB control number for this project is 0915-0327. Public reporting burden for this collection of information is estimated to average .5 hours per response, including the time for reviewing instructions, searching existing data sources, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to HRSA Reports Clearance Officer, 5600 Fishers Lane, Room 14N136B, Rockville, Maryland, 20857.

