## Negotiation Data Elements ICR Crosswalk of Changes Between 60-Day Notice and 30-Day Notice

Location of Edits <sup>i</sup>	Summary of Changes (Following 60-day Comment Period)	Type of Change	Explanation of Changes
Supporting Statement (throughout), Negotiation Data Elements ICR Form (hereinafter, "ICR Form" (throughout))	<ul> <li>Revisions to terminology, for consistency with the revised program guidance issued concurrently with the 30-day comment period, including:         <ul> <li>Section B: Clarified definition of non-FAMP and added definition of non-FAMP dosage form unit;</li> <li>Section C, Question 2 definition: Clarified that the basic pre-clinical research period is determined in reference to each indication of the selected drug;</li> <li>Section C, Question 3 definition: Broaden and clarified direct costs that should be included in post-IND costs;</li> <li>Section C, Question 4 definition: Added definition of therapeutic class for purposes of reporting costs of failed or abandoned products</li> </ul> </li> </ul>	Modify / Add	Technical changes to be consistent with policy changes in response to comments and / or due to internal or administrative review, including guidance review

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	related to the selected drug.  Section C, Question 6 definition: Moved definitions of U.S. net revenue from Section G; added questions, instructions, and definitions for 6a and 6b.  Section F: Modified language in patent and exclusivity definitions and clarified types of FDA applications and approvals about which CMS is seeking information;  Section G: Clarified patient assistance programs to explain definition; removed certain definitions (e.g., 340B ceiling price, 340B prime vendor program price, covered entity, quarterly total U.S. unit volume, manufacturer average net unit price);		

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	and removed definition of gross revenue; Section G: Added footnote to Big Four price definition; Section G, Question 16: Changed reporting for WAC from NDC-9 to NDC- 11; Section G, Questions 24- 25: Changed reporting for U.S. commercial average net unit price from NDC-9 to NDC-11; Section H, Question 28: Added definition of therapeutic alternative; Section H, Question 29: Added definitions of therapeutic alternative, outcomes, and patient- centered outcome; Section H, Question 30: Added definition of specific populations and health equity; and Section H, Question 31: Revised definition of unmet medical need.		

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Supporting Statement (Justification)	<ul> <li>Moved location of information required by the Primary Manufacturer, added data factors CMS will consider in negotiating a maximum fair price and removed inapplicable examples, revised language to be consistent with guidance revisions</li> <li>Updated language regarding submission methods to be consistent with guidance revisions</li> </ul>	Modify / Add	Changes due to internal or administrative review, including guidance review
Supporting Statement (Federal Register)	<ul> <li>Revised language consistent with publishing a revised package for a 30-day public comment period and for added specificity</li> </ul>	Add	Technical update
Supporting Statement (Cost to the Federal Government, Changes to Burden)	<ul> <li>Revised federal burden estimate to include the cost for modifications to the CMS HPMS system to include this ICR</li> </ul>	Modify	Changes due to internal or administrative review
ICR Form (throughout)	Revised section lettering and question numbering	Modify	Technical changes to be consistent with policy changes in response to comments

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ICR Form (General Instructions)	<ul> <li>Revised formatting of sections and language to clarify submission method for the public</li> </ul>	Modify	Technical update
ICR Form (General Instructions)	<ul> <li>Removed references to questions removed from this ICR in Section G</li> </ul>	Modify	Policy change in response to comments
ICR Form (General Instructions)	<ul> <li>Modified language about calculating and reporting monetary values consistent with the ICR and generally accepted accounting principles</li> </ul>	Modify	Technical changes to be consistent with guidance and external review
ICR Form (General Instructions)	<ul> <li>Revised instructions for Primary Manufacturers to not adjust any reported dollar amounts for inflation</li> </ul>	Modify	Policy change in response to comments
ICR Form (Section A, Instructions and Questions)	<ul> <li>Revised instructions and questions to explain that CMS will pre-populate certain information for NDC-11s and ask Primary Manufacturers to supplement as applicable</li> </ul>	Modify / Add	Technical changes to be consistent with guidance and external review
ICR Form (Section B, Instructions and Question)	<ul> <li>Revised instructions and table to include submission of dosage form unit</li> </ul>	Modify	Policy change in response to comments
ICR Form (Section C, Instructions)	<ul> <li>Revised introductory language and instructions to match revisions made to questions</li> </ul>	Modify	Policy change in response to comments

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ICR Form (Section C, Question 1)	<ul> <li>Moved and modified definition, questions, and instructions for reporting of acquisition costs from Section G to new question 1</li> </ul>	Modify/Revise	Policy change in response to comments
ICR Form (Section C, Questions 3 and5)	Revised "Post-Investigational New Drug (IND) Application Costs" to include "Completed U.S. FDA-Required Post-marketing Trials" and "Other R&D Direct Costs" to include other post-marketing trials and clarified how "all other R&D direct costs" should be reported	Modify	Policy change in response to comments
ICR Form (Sections C and E, Questions 3, 5, and 10)	<ul> <li>Increased word limit because multiple questions were combined</li> </ul>	Modify	Technical update
ICR Form (Section C, Question 5)	<ul> <li>Added definition and instructions</li> </ul>	Modify	Change due to internal or administrative review
ICR Form (Section C, Questions 6, 6a, and 6b)	<ul> <li>Split question 6 into two components: 6a and 6b</li> <li>Revised to add information previously in Section G to questions 6, 6a and 6b regarding global and U.S. net revenue metrics</li> <li>Revised reporting of U.S. net revenue from quarterly to lifetime</li> </ul>	Modify / Add	Policy change in response to comments

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ICR Form (Section D)	<ul> <li>Revised manufacturer submission of production and distribution costs from NDC-9 to NDC-11 (CMS will aggregate multiple NDC-11s to NDC-9s)</li> </ul>	Modify	Policy change in response to comments
ICR Form (Section D)	<ul> <li>Added description of "labeling"</li> </ul>	Add	Changes due to internal or administrative review
ICR Form (Section E, Instructions)	<ul> <li>Revised to clarify how indirect costs should be considered</li> <li>Revised to describe how responses to questions 9-11 should be calculated</li> </ul>	Modify / Add	Changes due to internal or administrative review
ICR Form (Section E, Questions 9, 10)	<ul> <li>Revised language to include all funding in question 9 and to specify sources of the funding in question 10</li> </ul>	Modify	Policy change in response to comments
ICR Form (Section F, Instructions)	<ul> <li>Revised description of relevant time period for reporting patent information</li> </ul>	Modify	Policy change in response to comments
ICR Form (Section F, Question 12)	<ul> <li>Revised example language to illustrate the types of patents and patent applications about which CMS seeks information and modified response format for pending patent applications</li> </ul>	Modify/Add	Policy change in response to comments

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ICR Form (Section F, Question 13)	<ul> <li>Revised language to clarify that manufacturers should describe patents for the selected drug that are listed in the FDA Orange or Purple Book</li> </ul>	Modify	Technical change in response to comments
ICR Form (Section F, Question 12)	<ul> <li>Added example for submission of information on the purpose of a patent</li> </ul>	Modify/Add	Technical change in response to comments
ICR Form (Section F, Question 14)	<ul> <li>Revised language to clarify that exclusivities are statutory and that FDA has not determined the first licensure for each 351(a)- biological product in the Purple Book</li> </ul>	Modify	Technical change in response to comments
ICR Form (Section F, Question 15)	<ul> <li>Added language for manufacturers to submit any efficacy supplements that have been approved or are pending FDA approval</li> </ul>	Modify/Add	Policy change in response to comments
ICR Form (Section F, Question 15)	<ul> <li>Removed column for manufacturers to enter submission number and modified response format for application status and classification code</li> </ul>	Modify	Technical change in response to comments; Change due to internal or administrative review
ICR Form (Sections H-J	<ul> <li>Moved Certification of Submission of Sections A through G (formerly Section I) to follow section G; moved</li> </ul>	Modify	Technical changes due to internal or administrative review

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	Evidence on Alternative Treatments (formerly Section H) to section I		
ICR Form (Section H)	<ul> <li>Revised language to clarify which individuals of the Primary Manufacturer are eligible to certify the data submission</li> </ul>	Modify	Changes due to internal or administrative review
ICR Form (Section I, Question 26)	<ul> <li>Revised language to capture additional types of respondents, including researchers and caregivers</li> </ul>	Modify	Technical change in response to comments
ICR Form (Section I, Instructions for Questions 27-31)	<ul> <li>Revised language to clarify evidence and documentation requirements if evidence includes or is related to qualify-adjusted-life-years (QALYs)</li> <li>Added language to clarify evidence and documentation requirements if evidence is related to cost-effective measures</li> <li>Clarified instructions regarding use of citations</li> <li>Added instructions pertaining to caregivers who may respond to questions in Section H</li> </ul>	Modify / Add	Policy change in response to comments

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	<ul> <li>Added instructions regarding the permissible submission of visual representations</li> </ul>		
ICR Form (Section I, Instructions for Questions 27-31)	<ul> <li>Removed duplicate bullet regarding instructions for submission of materials</li> <li>Added clarification of question instructions for unpublished material, citations and visual representations of information</li> </ul>	Modify	Technical change due to internal review
ICR Form (Section I, Question 27)	<ul> <li>Added clarification of question instructions for off- label uses</li> <li>Removed submission of FDA labels, as CMS is able to view this information</li> </ul>	Add / Modify	Policy change in response to comments  Technical change due to internal review
ICR Form (Section I, Questions 28-30)	<ul> <li>Added ability to submit visual representation of information (e.g., charts/tables/graphs)</li> </ul>	Add / Modify	Policy change in response to comments
ICR Form (Section I, Questions 27-30)	<ul> <li>Added language to checkbox clarifying that the response also includes measures used in the evidence submitted</li> </ul>	Add	Changes due to internal or administrative review
ICR Form (Section I, Question 28)	<ul> <li>Clarified language regarding outcome information requested</li> </ul>	Modify / Add	Policy change in response to comments

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	<ul> <li>Added language requesting key outcomes and explanation of selection</li> <li>Added language regarding risks, harms, side effects or unique scenarios or considerations of the drug or therapeutic alternative</li> </ul>		
ICR Form (Section I, Question 29)	<ul> <li>Revised to clarify individuals who are terminally ill and request explanation of specific populations noted</li> <li>Added question regarding health equity considerations</li> </ul>	Modify / Add	Policy change in response to comments
ICR Form (Section I, Question 30)	<ul> <li>Added a question regarding unmet medical need</li> </ul>	Add	Policy change in response to comments
ICR Form (Section I, Question 31)	<ul> <li>Added a new question and corresponding instructions to capture patient and caregiver experiences related to the selected drug and/or therapeutic alternatives</li> </ul>	Add	Policy change in response to comments
ICR Form (Section I, Question 32)	Added a new question (required for manufacturers; optional for the public— except not permitted for patients/caregivers) for submission of an executive summary of information provided in response to questions 27-30	Add	Policy change in response to comments

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ICR Form (Section J)	<ul> <li>Revised certification language for submissions of data in response to Section I</li> </ul>	Modify	Policy change in response to comments

<sup>&</sup>lt;sup>i</sup> References to section and question numbers reflect the lettering and numbering in the revised 30-day notice.