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EPD use levels are cumulative. Transparency is the baseline. To create a 'Data source' conformant PCR, all criteria in all checklists must be documented.

1. Program Operator (PO) checklist Version 1.0, May 25, 2022 | ACLCA PCR Open Standard 2022

| | | | | | | | 3 Data source | |
|----------------|-----|-------|---|---|--|----------------|---|----------|
| Categories | | # | Criteria | ISO reference | Supporting documentation | EPD use | 2 Procurement | |
| | | | | | | | 1 Transparency | |
| | Gro | und r | | | | | How criteria were met | Due |
| Organizational | Ø | | Prior to using the ACLCA PCR Guidance 2022 to develop PCRs, the PO shall use this guidance to develop and publish conformant program instructions that describe the process of PCR development aligned with ISO/TS 14027. | This guidance | General program instructions (governance document): • ACLCA PCR Guidance 2022 conformant statement with version number | 1 Transparency | Updated program instructions published to SM website http://www.sustainableminds.com/files/transparency/SM_Governance _and_program_rules.pdf | Complete |
| | ☑ | 2 | PO shall use this checklist to guide the creation of a PCR, identify how criteria were met, and provide the completed Program Operator Checklist and PCR Review Panel Checklist to the PCR Review Panel. | This guidance | PCR supporting documentation: • Completed checklist | 1 Transparency | Completed checklists saved with the PCR supporting documentation. | Complete |
| | | | PO shall be the secretariat of the PCR and manage an open and transparent process to develop or update a PCR. This process shall include public notices prior to PCR development and an open consultation process with interested parties while the PCR Committee remains active. | | | | Public notice on the Sustainable Minds website announcing the new lavatories Part B on June 11, 2024: http://www.usstainableminds.com/transparency-report-program/part- | |
| | ☑ | 3 | PO shall publish the intention to develop (or update) a PCR on its website, in relevant industry and trade publications and/or news services, and through centralized notification mechanisms. The announcements shall include contact information that allows interested parties to request more information about participation in the PCR development or review process. | 14027 Clause 6.4.1 | PCR supporting documentation: • Date(s) announcement(s) were posted and where | 1 Transparency | b Email blast on June 11, 2024 to mailing lists of LCA professionals, building and construction industry and trade associations, and manufacturers with published transparency documentation listed in the Transparency Catalog under the plumbing CSI MasterFormat Division (22 00 00). | Complete |
| | | | Interested parties may include material suppliers, manufacturers, trade associations, purchasers (such as architects, designers, specifiers, contractors, and engineers), users, non-governmental organizations (NGOs), and public agencies. | | | | (22 00 00). | |
| | Ø | 4 | PO shall determine whether to create a new PCR or to adapt an existing PCR from other geographic regions. The PO shall justify the determination in the PCR. | 14027 Clause 6.4.2, 6.4.3 | PCR: • Identify existing PCRs considered, and provide justification for creating a new PCR. • If new, identify the supporting LCA. • Describe how existing PCRs will be adapted. | 2 Procurement | N/A | N/A |
| | ☑ | 5 | PO shall evaluate upstream and downstream PCRs in the value chain to be considered for alignment. PO shall list relevant PCRs in the PCR. Note: Also see Criterion 15 for the process of determining when a PCR may be updated. | 14044 14027 Clause 6.4.3 This guidance | PCR supporting documentation: • Identify existing upstream PCRs for the major inputs to the product(s) considered in the PCR. • Describe differences in allocation rules or other potential conflicts and how they were resolved. • Identify existing downstream PCRs that use products/materials from the PCR and how inconsistencies were resolved. | 3 Data source | N/A | N/A |
| | Ø | 6 | PO shall harmonize PCR activities with other EPD programs to avoid unnecessary duplication and proliferation of similar PCRs, and align on mutual recognition agreement (MRA) requirements. PO shall list relevant PCRs in the PCR. Note: Refer to both the ACLCA's PCR library and the North American PCR Catalog: Building & Construction Materials https://www.transparencycatalog.com/na-pcr-catalog-building-products | 14029 Clause 7, 9.2 | PCR supporting documentation: • Identify whether this criteria is applicable. • Identify other POs engaged to harmonize PCR activities and opportunities explored (joint development of new, merging, application of existing, or adaption of existing). • MRA between POs one exists. | 1 Transparency | Addressed in Program operator responsibilities section of Part B. | Complete |
| | Ø | 7 | PO shall publish and implement procedures for an appeals mechanism to ensure prompt and impartial handling of procedural complaints regarding any action or inaction of the PCR Committee, PCR Review Panel, or Program Operator. | 14027 Clause 6.4.4 | General program instructions (governance document): • Explanation of appeals process | 1 Transparency | Addressed in section 10.0 of the governance document. | Complete |
| | Ø | 8 | PO should include a method for addressing data quality in its general program instructions. Note: Refer to the addendum "Assessing Data Quality of Background Life Cycle Inventory Datasets" for an example data quality assessment method. | | General program instructions (governance document): - Method for Data Quality Assessment | 2 Procurement | N/A | N/A |
| | PCF | R com | mittee formation | | | | How criteria were met | Due |
| | ☑ | 9 | PO shall actively reach out to interested parties (including parties outside the PO's country or region) to ensure that the PCR Committee is composed of independent members, making sure that the interests of one party do not dominate the PCR development process. No single interested party category (at individual, organizational, or sectoral levels) shall dominate the membership of a PCR Committee. Interested parties may include material suppliers, manufacturers, trade associations, purchasers (such as architects, designers, specifiers, contractors, and engineers), users, non-governmental organizations (NGOs), and public agencies. | 14025 Clause 5.5, 6.5, & 9.3 14027 Clause 6.4.1 and 6.4.2 | PCR: • List of PCR Committee members with employer and/or other entity on behalf of which they are participating. PCR supporting documentation: • Description of interested party outreach efforts and explanation of interested party outreach. | 1 Transparency | Working group members listed on page 1 of each Part B. | Complete |

| | ☑ <i>'</i> | 10 | PO shall address potential conflicts of interest developing the PCR and fully disclose funding sources for the management to interested parties. If significant external funding was made by one or more parties to support the development, the PO should put in place procedures to ensure that no conflict of interest occurrs in the PCR process. 'Significant funding' is defined as more than \$10,000 or its in-kind equivalent, or 20% or more of the anticipated funding needs. | Performance Standards and Ecolabels for Federal Purchasing. | PCR supporing documentation: - The policy or procedure in use when the PCR was developed covering conflicts of interest, separation of organizational functions necessary to address any potential conflict of interest. - Attestation that this policy or procedure was followed during the development. The evidence must also include one of the following: - Documentation that original sources of funding were disclosed to interested parties, such as a disclosure statement, or in meeting minutes for relevant working groups. | 1 Transparency | Conflict statement included in the Part B development information table of each Part B. | Complete |
|---|------------|------|---|--|--|----------------|---|----------|
| (| onter | nt o | f PCR | | | | How criteria were met | Due |
| | ☑ . | 11 | The PCR shall report on the following items: Name and registration number of the PCR General information about the program: name of the program, contact information, ogo, and website if applicable PCR Committee members and affiliations Publication date Expiration date and renewal schedule Types of product claims covered by the PCR, with references to standards Product category Geographical representativeness of the PCR Original language and translations (if existing) How to make comments to the PCR | 14027 Clause 6.5 | PCR: • Draft PCR that includes all items reported | 1 Transparency | Part A section 1.1 addresses the use of SM PCRs to create ISO 14025 Type III environmental declarations, and also language availability. http://www.sustainableminds.com/files/transparency/SM_Part_A_LCA_ calculation_rules_and_report_requirements_2023.pdf All other items are addressed in each Part B. | Complete |
| | ☑ . | 12 | The PCR shall report the following information about the review process and background of the PCR: Review panel member information Open consultation period and participants Other existing PCRs for the product category and reasons for developing a new one Reference to underlying LCAs Confirmation statement that the PCR was created in conformance with this ACLCA PCR Guidance (including version number) | 14025 Clause 5.5, 8.2 14027 Clause 5.2, 6.4.4 14025 Clause 6.7.1, 6.7.2 14027 Clause 6.1, 6.4.3, 6.5.3, 7.1d | PCR: • Draft PCR that includes all items except 'open consultation period' PCR supporting documentation: • Open consultation period and participants | 1 Transparency | All items except open consultation participants addressed in Part B. Aggregated technical and public comments spreadsheet, including commenter names and committee responses, to be created and made available in the Detailed Review Report. | Complete |
| | CR re | evie | w process | | | | How criteria were met | Due |
| ı | | 13 | PO shall set up an independent third-party review panel composed of a minimum of hree members (a chair and two members). The combined competencies of the panel | 14027 Clause 7.1, 7.2, 7.3, 14025 Clause 8.2.3 | PCR: List of review panel members | 1 Transparency | Working group members listed on page 1 of each Part B. | Complete |
| | Ø , | 14 | PO shall also set up an open consultation review. | 14027 Clause 6.4.4, 7.3 | PCR supporting documentation: • Date(s) open consultation period(s) announced, where/how; aggregated comments spreadsheet | 1 Transparency | Aggregated technical and public comments spreadsheet, including commenter names and committee responses, to be created and made available in the Detailed Review Report. | Complete |
| | ⊻ | | PO shall ensure the PCR Review Panel provides comments within a 90-day period. | This guidance | PCR supporting documentation: • Date(s) PCR review period | 1 Transparency | Due date less than 90 days provided to PCR reviewer (Jul 17 - Aug 2). | Complete |
| | ublic | | n, new and updated PCRs | | | | How criteria were met | Due |
| | | 16 | he schedule for renewal, if applicable. | 14025 Clause 6.4, 6.7.1 14027 Clause 8.1.1 This guidance | PCR supporting documentation: • URL of PO's published PCRs page • URL PCR will be available at when published PCR: • Validity period of PCR • Conformance statement and EPD use case level | 1 Transparency | A link to the SM Part B page is included in each Part B. Completed Part Bs will be uploaded to that page when published. The URL of the Part B when published will be as follows: http://www.sustainableminds.com/files/transparency/pgds/Part_B_Product_Group_Definition_Commercial_Lavatories_2024.pdf Part B contains validity period, conformance statement, and EPD use case level. | Complete |

| | V | | To manage the expectations of PCR users, the PO shall post update information on its website at least four months in advance of the expiration date. The update options include: extending the current PCR, updating the PCR, or letting the PCR expire with no update. If information is not provided within this timeframe, other POs may proceed with the update and post PCR update information on their website. | This guidance | • URL of PO's PCRs undergoing updates | 1 Transparency | Public notice on the Sustainable Minds website announcing the new lavatories Part B on June 11, 2024: http://www.sustainableminds.com/transparency-report-program/part-b Email blast on June 11, 2024 to mailing lists of LCA professionals, building and construction industry and trade associations, and manufacturers with published transparency documentation listed in the Transparency Catalog under the plumbing CSI MasterFormat Division (22 00 00). | Complete |
|----------------|-----|-----|---|--------------------|--|----------------|---|----------|
| | Ø | | To update a PCR during the validity period, the PO shall: 1. Notify the original PCR Committee members and original Review Panel. 2. Consult ISO 14027 to confirm the reason to update is valid. 3. Create or update the ACLCA PCR Guidance Checklists for the PCR. 4. Open consultation to interested parties. 5. Update the PCR. 6. Obtain sign-off by PCR Review Panel. 7. Republish an updated version and include a change log at the start of the document. 8. Announce the updated version. 9. Update the ACLCA PCR Repository. In the case that an existing PCR does not meet the requirements for creating EPDs for public or private procurement purposes, the PO shall make an effort to first engage the commissioner of the PCR to reconvene the PCR Committee in order to make the required updates. If the PCR commissioner does not reconvene the PCR Committee within 30 days of the PO's request, then the PO may proceed to develop a new PCR using the existing PCR as an informative input document. | 14027 Clause 9 | PCR: • Valid update reason PCR supporting documentation: • Checklists | 1 Transparency | The Part B development information table in each Part B lists an Update justification where relevant. For this Part B, updates were not made during the validity period. The process for updating a PCR during the validity period is included in section 9.0 of the governance document. http://www.sustainableminds.com/files/transparency/SM_Governance_and_program_rules.pdf | Complete |
| | Ø | 19 | For substantial PCR updates (e.g., updates that impact the results of an EPD), the PO shall contact manufacturers in their program with valid EPDs and other POs to bring attention to the PCR changes and encourage that they update accordingly. | 14027 Clause 9 | PCR supporting documentation: Description of notification and dates of outreach | 1 Transparency | Not applicable; this is a new Part B. | Complete |
| | EPD | tem | plate | | | | How criteria were met | Due |
| | Ø | 20 | PO shall create a standard EPD template to be used for all EPDs that can be customized per PCR to identify requirements unique to each. Consider both digital and print (PDF) publishing. Note: Refer to the 'EPD Comparatibility and Digital EPDs / Open EPD addendum. PO shall include a statement adjacent to the PCR name to indicate conformance with this guidance and the EPD use case level. | This guidance | PCR: • EPD template document prepared for this PCR • Statement text included in EPD template | 1 Transparency | A standard EPD template is included in Appendix C of Part A. Under the name of each Part B is a statement indicating conformance to this guidance and the EPD use case level. | Complete |
| | Ø | 21 | PO shall ensure that the type of EPD developed is clearly noted on the EPD. <i>Note:</i> Refer the 'EPD Types' addendum. | This guidance | PCR: • Statement text included in EPD template | 1 Transparency | Requirement listed in the Verification statement section in Appendix C of Part A (EPD template). | Complete |
| Goal and scope | ☑ | 22 | Product categories shall be primarily defined and sufficiently described by product functionality, technical performance, and use. The PCR shall clearly define the product groups for which the rules apply, both by using descriptive language and by using the relevant codes for any of the existing classification systems relevant to the product category and region. Products NOT covered by the PCR shall be clearly listed (as a clarification when products are similar). PO should ensure that the product classification systems are not to be the single determining factor for defining the product category. The PCR is encouraged to provide sufficient information to clearly describe the scope of products and services for which the rules apply. | 14027 Clause 8.1.1 | PCR: • Draft PCR which includes all the items | 2 Procurement | N/A | N/A |

EPD use levels are cumulative. Transparency is the baseline. To create a 'Data source' conformant PCR, all criteria in all checklists must be documented.

2. PCR Committee checklist Version 1.0, May 25, 2022 | ACLCA PCR Open Standard 2022

October 2, 2024 | Sustainable Minds | Contact Kim Hammer (kim@sustainableminds.com)

| Categories | # | Criteria | ISO reference | Supporting documentation | EPD use | 3 Data source 2 Procurement 1 Transparency | |
|----------------|-------------|---|--|--|----------------|--|----------|
| Description | Ground | ulae | | | | How criteria were met | Due |
| Documentation | | PCR Committee shall use this checklist to guide the creation of a PCR, identify how criteria were met, and provide the completed checklist to the Program Operator to provide to the PCR Review Panel. | This guidance | PCR supporting documentation: • Completed checklist | 1 Transparency | Completed checklists saved with the PCR supporting documentation. | Complete |
| | ☑ 2 | PCR Committee shall thoroughly document the use of an existing PCR as an informative document in any adaptation to create a new PCR. Include the PO name, existing PCR name, product category classification, link to the existing PCR, and provide justification for adapting the existing PCR. | 14027 Clause 6.4.3 and this guidance | PCR: • Link to PCR Committee's documentation of adaptation | 2 Procurement | N/A | N/A |
| | ☑ 3 | PCR Committee shall respond to each comment from the PCR Review Panel and public consultation. Responses should address any conflicting comments provided by the PCR Review Panel. | This guidance | PCR supporting documentation: • Link to PCR Committee's documented public response to comments and consultation on PO's website (aggregated comments spreadsheet). | 1 Transparency | Aggregated public comments and review panel comments, including committee responses, created and published on the SM website with the PCR supporting documentation. | Complete |
| | ✓ 4 | PCR Committee shall provide a limited description of the involvement of interested parties for open consultation. Specifically, the PCR should provide: • The name and/or affiliation of the stakeholders who participated in the open consultation. • The dates of the open consultation period. Public consultation should be utilized during the PCR review process. The public consultation of the completed draft PCR should include at a minimum a 30-calendar-day time period for comments to be submitted. | 14025 Clause 5.5 14027 Clause 5.2, 6.4.4 | PCR: • Draft PCR that includes list of participating interested parties and dates of consultation period. | 1 Transparency | Open consultation period listed in 'Open consultation' section of the Part B development table. Aggregated technical and public comments spreadsheet, including commenter names and committee responses, to be created and made available in the Detailed Review Report. | Complete |
| Compliance | ☑ 5 | PCR Committee shall ensure that the underlying LCA meets the requirements of ISC 14044 and other pertinent standards and that, according to these standards, it has either been critically reviewed by a third party or has undergone an internal verification, either by the PCR Committee itself or appointed independent LCA expert. | 14025 Clause 6.7.1, 6.7.2, 8.1.3, 8.2.1, 8.2.2 | PCR supporting documentation: Link to documentation of LCA review or internal verification. | 2 Procurement | N/A | N/A |
| | ☑ 6 | PCR Committee shall ensure that the PCR is compliant with any referenced standards and relevant program instructions under which it is developed. | | PCR: • List of referenced standards and link to relevant program instructions. | 1 Transparency | Use of Part B in conjunction with SM Part A is addressed in Program operator responsibilities section of each Part B. SM Part A section 1.1. lists the standards required for conformance. The last section of each Part B contains a link to where to find the SM program instructions (governance document). | Complete |
| □ 7 | | PCR Committee shall establish LCA requirements that are consistent with ISO 14044. The PCR Committee is encouraged to develop end-use case scenarios for the PCR-compliant EPDs and to incorporate considerations for these use cases into the underlying LCA. | 14025 Clause 6.7.1, 6.7.2 14027 Clause 5.1, 6.1, 6.5.3, 7.1d | PCR supporting documentation: Third-party reviewed ISO 14040/44 conformant LCA of the product categories under consideration. The LCA will reflect cases in which the EPD may be interpreted in use. | 1 Transparency | A link to the underlying LCA is included in the Program operator responsibilities section of Part B. | Complete |
| | Ground | ules | | | | How criteria were met | Due |
| Goal and scope | | PCR Committee shall ensure that all rules for LCA are specified and harmonized wit upstream and downstream PCRs (if available) in conformance with relevant standards, including: specification of the functional unit, scope of the study, inventor collection, any allocation rules, impact assessment, and rules for additional information. | 14044 | PCR: • Draft PCR with list of specifications | 3 Data source | N/A | N/A |
| | ☑ 9 | PCR Committee shall ensure that the product category used in the underlying LCA supporting the PCR is directly applicable to the PCR. | 14025 Clause 3.14, 6.6, 6.7.2 14027 Clause 6.5.2, 6.5.3 | PCR: • Specification and justification of the product category and applicable functional unit. | 2 Procurement | N/A | N/A |
| | ☑ 10 | PCR Committee shall define the study scope and EPD type for construction product and services. | 21930 Clause 5.2.1, 5.2.2 | PCR: • Draft PCR with specification of scope as cradle-to-gate or cradle-to-gate with options or cradle-to-grave. | 1 Transparency | Part B specifies the scope as as cradle-to-grave. | Complete |
| | ☑ 11 | PCR Committee shall ensure that a clearly defined and measurable functional or declared unit is included in the PCR for construction products and services. | 21930 Clause 7.1.2, 7.1.3 | PCR: • Draft PCR with detailed description of the application and suitability of defining functional and declared units, respectively. | 1 Transparency | Part B provides a description of the functional unit. | Complete |

| PCC Committee shall determine the level of gramwardy of unit processes specified by the PCR to be included in the curelings (CA apportune the SPD and entire the PCR supprise) and the supprise of the supprise that the supprise | Ø | 12 | The PCR Committee shall determine which EPD types may be developed (ex: product-specific, industry-wide) and state the specific data requirements for each type. Any other terminology describing types of EPDs should be discouraged. <i>Note: Refer to the 'EPD Types' addendum for descriptions</i> . | ISO 21930 Annex B and 'EPD Types' addendum | PCR: • Draft PCR with description of the EPD types with specific data requirements | 1 Transparency | Part B specifies EPD type under the name of the Part B. Specific data requirements are listed in the Additional rules to Part A section of Part B. | Complete |
|--|------|--------|--|---|---|----------------|---|----------|
| PCS Committee deal forces part of any object of production of the | Sys | tem b | oundary | | | | How criteria were met | Due |
| 1) of minimum, according togother of uniformity and many of white the control of the policy of the p | | | PCR Committee shall determine the level of granularity of unit processes specified by the PCR to be included in the underlying LCA supporting the EPD and ensure that these are consistent with the study's goal of using well-identified and explained | 14027 Clause 6.5.3 21930 Clause 7.1.9 for construction products & | Draft PCR with list of all unit processes that include all service, material, and energy flows directly connected to the study project and | 3 Data source | N/A | N/A |
| infrastructure to be included in cases wherever it is featable. The PCR Committee is the contracting of pages by the interest or committee in the contracting pages and in the position of the | ☑ | 14 | at minimum, a cradle-to-gate[1] system boundary and that any deviation is explicitly specified and justified; and 2) the use of the recycled content (i.e., cut-off) approach for end-of-life allocation of environmental burdens between product systems. 1] "Gate" represents the finished and packaged product at the manufacturing facility just prior to | 14025 6.7.2b, 6.7.2c, 6.7.2j, 7.2.5 | Draft specification of the system boundary and justification of any | 2 Procurement | N/A | N/A |
| standard guidelines for any and each life cycle stage to be included beyond cradle-to-gate (i.e., A.1-A.5) the PCR sogne and require LCA results for these be reported. This guidance PCR: Where applicable, list of scenarios and associated assumptions. PCR committee shall specify whether the benefits and loads beyond the system boundary (i.e., Module D) are to be included in the PCR of red eviation from the prescribed scenarios. PCR Committee shall specify whether the benefits and loads beyond the system boundary (i.e., Module D) are to be included in the PCR of the stage of the specified scenarios. PCR Committee shall specify whether the benefits and loads beyond the system boundary (i.e., Module D) are to be included in the PCR of the specified and reported in practice of a product. Note: Refer to the "Circular Scenarios (Module D)" addendum This guidance and Circular Scenarios (Senarios) senarios (Senarios (Module D)" addendum senarios (Module O)" addendum senarios (Module | Ø | 15 | infrastructure to be included in cases whenever it is feasible. The PCR Committee is encouraged to specify lifetimes or standardized methods of computing lifetimes, as well as the depreciation method utilized to allocate the burden of capital goods over their service period, with any deviations from the default approach explicitly specified | This guidance | | 2 Procurement | N/A | N/A |
| boundary (i.e., Module C) are to be included in the EPD. Iso, the PCR shall reported to separately in relevant EPDs communicating the full life cycle (cradie-to-grave) impacts of a product. Note: Refer to the 'Circular Scenarios (Module D)' addendum. Data collection | Ø | 16 | standard guidelines for any and each life cycle stage to be included beyond cradle-to- gate (i.e., A1-A3) in the PCR scope and require LCA results for these be reported. The PCR shall also prescribe assumptions for scenarios in cases where there is no discernable difference between one product and another in the same category for use and end-of-life stages. The PCR Committee should include criteria in the PCR | | | 2 Procurement | N/A | N/A |
| Life cycle inventory PCR Committee shall prescribe acceptable primary data collection practices and clearly specify the scope and data quality for secondary data with recommendations for use of specific datasets or databases facilitating this process. Datasets used for calculations shall have been updated within the last 10 years for producer-specific (foreground) data; deviations shall be proposed for geographic areas or technologies beyond the scope of the specified dataset(s). Any deviation from the recommended background (secondary) datasets in the PCR shall equire EPDs to disclose the reporting period for primary and secondary data. Nate: Refer to the Assessing Data Quality of Background Life Cycle Inventory Datasets' addendum. PCR: **Oratif PCR that includes all items** **PCR: **Oratif PCR that includes all items** **PCR: **Draft PCR that includes all items** **PCR: **Draft PCR that includes all items** **PCR: **Draft PCR that includes all items** **I Transparency** **SM Part A includes the list of selected LCIA indicators. 6.6.6.6.6.6.6.6.6.6.6.6.6.6.6.6.6.6.6 | | | boundary (i.e., Module D) are to be included in the EPD. If so, the PCR shall describe the specific scenario(s), benefits, and loads to be considered and reported separately in relevant EPDs communicating the full life cycle (cradle-to-grave) impacts of a product. Note: Refer to the 'Circular Scenarios (Module D)' addendum. | 'Circular Scenarios (Module | Where applicable, list of scenarios and concomitant benefits and | 2 Procurement | n/A | N/A |
| PCR Committee shall prescribe acceptable primary data collection practices and clearly specify the scope and data quality for secondary data with recommendations for use of specific datasets or databases facilitating this process. Datasets used for calculations shall have been updated within the last 10 years for background data and within the last 5 years for producer-specific (foreground) data; deviations shall be justified. Where databases are required, alternatives or modifications shall be proposed for geographic areas or technologies beyond the scope of the specified dataset(s), Any deviation from the recommended background (secondary) datasets in the PCR shall be clearly specified and justified. In addition, the PCR shall require EPDs to disclose the reporting period for primary and secondary Datasets' addendum. PCR: * Draft PCR that includes all items * Dra | Data | a coll | ection | | | | How criteria were met | Due |
| 19 indicators or additional information requirements for which relevant inventory information shall be collected. 1 Transparency SM Part A includes the list of selected LCIA indicators. 1 Transparency SM Part A includes the list of selected LCIA indicators. | Ø | | clearly specify the scope and data quality for secondary data with recommendations for use of specific datasets or databases facilitating this process. Datasets used for calculations shall have been updated within the last 10 years for background data and within the last 5 years for producer-specific (foreground) data; deviations shall be justified. Where databases are required, alternatives or modifications shall be proposed for geographic areas or technologies beyond the scope of the specified dataset(s). Any deviation from the recommended background (secondary) datasets in the PCR shall be clearly specified and justified. In addition, the PCR shall require EPDs to disclose the reporting period for primary and secondary data. Note: Refer to the 'Assessing | 'Data Quality and Secondary Background | PCR: | 2 Procurement | n/a | N/A |
| | Ø | 19 | indicators or additional information requirements for which relevant inventory | 14027 Clause 6.5.4, 6.5.5, | | 1 Transparency | SM Part A includes the list of selected LCIA indicators. | Complete |
| PCR Committee shall specify, based on the underlying LCA and/or additional studies 14025 Clause 7.2.3, 7.2.4 informing the PCR, all the data that are to be collected (rather than specifying cut-off 14027 Clause 6.6 Procurement 14027 Clause 6.6 Proc | Ø | 20 | informing the PCR, all the data that are to be collected (rather than specifying cut-off | 14025 Clause 7.2.5, 7.2.4 | | 2 Procurement | N/A | N/A |
| PCR Committee shall specify the type of data to be collected. The committee is encouraged to follow standard data collection examples for foreground (primary) data 21930 Clause 7.1.9 and 14044 Annex A 21930 Clause 7.1.9 braft PCR with data collection sheet example specific to PCR 2 Procurement N/A | | | encouraged to follow standard data collection examples for foreground (primary) data collection. | | 7 7 11 | 2 Procurement | n/A | N/A |
| Data quality How criteria were met | Data | a qua | lity | | | | How criteria were met | Due |

| | | PCR Committee shall refer to relevant guidance to consider parameters for assessing data quality of both foreground (primary) and background (secondary) data. Note: Refer to the 'Assessing Data Quality of Background Life Cycle Inventory Datasets' addendum which provides a data quality assessment method. | 21930 Clause 7.1.9 14044 Clause 4.2.3.6 14025 Clause 6.7.2 14027 Clause 6.2 | PCR supporting documentation: • Complete data quality assessment for both foreground (primary) and background (secondary) data. This information shall also be included in the underlying LCA, and reviewed. | 1 Transparency | A data quality assessment of primary and secondary data is included in the underlying LCA and was reviewed by the PCR committee. | Complete |
|---|---------|--|---|---|----------------|--|----------|
| В | ckgrou | ind/secondary data | | | | How criteria were met | Due |
| | 23 | PCR Committee shall ensure that the PCR specifies background (secondary) data quality requirements such that differences between claim results are rooted in actual technical differences, rather than artifacts of background data or the platform. If a secondary data source does not meet the required quality specified by the PCR, it shall be verified by the program operator that better data is not available. Note: Refer to the 'Assessing Data Quality of Background Life Cycle Inventory Datasets' addendum which provides a data quality assessment method. For example, as detailed in this addendum, the most recent version of background data for baseline electricity from Federal LCA Commons met the data quality requirements and is recommended to be specified across PCRs (with the LCI and method compatible with the Federal Elementary Flow List (FEDEFL) from https://www.lcacommons.gov/. | Assessing Data Quality of Background Life Cycle Inventory Datasets' addendum | PCR: • Draft PCR with list of background (secondary) data sources and default LCIA method(s) | 2 Procurement | N/A | N/A |
| F | regrou | nd/primary data | | | | How criteria were met | Due |
| 1 | 3 24 | PCR Committee shall ensure that the PCR specifies primary data be collected for every process in the product system under the control of the organization making the product claim. The PCR Committee is encouraged to specify that data specific to the investigated product scope and supply chain are preferable to generic data, particularly in unit processes considered to have a significant contribution to the product life cycle. For EPDs seeking transparency-level conformance with this guidance, the PCR shall require the following: EPDs that use secondary data for any unit process that contributes 30% or more to any disclosed environmental impact category shall disclose the data source (database name and version, dataset name, dataset geography, and dataset allocation method). | This guidance | PCR supporting documentation: • Foreground (primary) data collected in conducting the underlying LCA, and the sensitivity of LCIA outcomes to variability in the foreground data. A facility-specific data collection protocol shall also be included. | 1 Transparency | SM Part A section 7.6 states that primary data shall be collected for every process in the product system under the control of the organization(s) developing the LCA. Part B contains a statement in the Additional rules to Part A section which states: EPDs that use secondary data for any unit process that contributes 5% or more to any disclosed environmental impact category shall disclose the data source (database name and version, dataset name, dataset geography, and dataset allocation method) The underlying LCA lists primary data collected and includes an analysis on sensitivity or variability. | Complete |
| | 1 25 | For EPDs seeking procurement-level conformance with this guidance, the PCR shall require that EPDs use facility-specific data for upstream unit processes that cumulatively contribute 50% or more to the disclosed global warming potential. In situations where facility-specific data is not available for the upstream unit processes, and such a facility is required to report to the EPA Greenhouse Gas Reporting Program (GHCRP), the PCR shall require the EPD to disclose in the Additional Environmental Information section: the carbon intensity of the manufacturing plant (carbon emitted per metric ton of product manufactured) from which these products, and/or the quartile in which in which the manufacturing plant resides where benchmarks have been published [https://www.epa.gov/ghgreporting/ghgrp-minerals]. Carbon intensity shall be calculated by dividing the emissions reported to the EPA GHGRP by plant production. Emission and production data must be from the same reporting period using the most recent year of data. When a published ENERGY STAR Energy Performance Indicator is available for a product or constituent upstream product, the PCR shall require the EPD to disclose in the Additional Environmental Information section: the ENERGY STAR Energy Performance Score for the manufacturing plant in which the product or constituent upstream product was manufactured, and the reporting period of the underlying data. See https://www.energystar.gov/industrial_plants/energy_star_plant_certification/buy_clean_procurement_and_energy_star_0 for more information. | This guidance | PCR: • Draft PCR that includes all items | 2 Procurement | N/A | N/A |
| | 26 | PCR Committee shall ensure that the PCR specifies the means by which primary data should be collected and may provide templates to facilitate harmonized data collection, metadata recording, and results reporting. If the specified data collection means are unachievable for a specific EPD developer, the PCR shall designate that the developer records the data collection method(s) utilized in the data description. | 14025 Clause 6.7.2 | PCR: • Specification of data collection methods (e.g., measured, calculated, estimated) | 1 Transparency | SM Part A section 7.6 states: The method of data collection shall be specified (e.g., measured, calculated, estimated). | Complete |

| | umptions | | | | How criteria were met | Due |
|-------------|--|---|---|----------------|---|----------|
| ☑ 27 | PUR Committee shall specify all parameters of assumed scenarios for use and end- of-life stages so as to ensure comparability and consistency of results. If a manufacturer wishes to define their own scenario(s), they shall be based on primary | This guidance and the 'Circular Scenarios (Module D)' and the 'Allocating Materials Shared Across Product Systems' addendu | PCR: • List of parameters for use and end-of-life stage scenarios | 2 Procurement | N/A | N/A |
| ☑ 28 | PCR Committee shall ensure that the PCR provides worst-case (i.e., 'conservative') default values for scenario data of the specified processes where no data are available for the EPD developer. | This guidance | PCR: • List of worst-case (i.e., 'conservative') default scenario values | 2 Procurement | N/A | N/A |
| Data com | npliance | | | | How criteria were met | Due |
| ☑ 29 | PCR Committee shall ensure that claims made in the PCR are based on the results of an LCIA, LCI, and/or substantiated and verifiable additional information modules relevant to the product category. | 14027 Clause 6.6 | PCR: • An underlying LCA with supporting LCIA and LCI for all PCR guidelines | 1 Transparency | The underlying LCA contains relevant supporting LCA results. | Complete |
| ☑ 30 | PCR Committee shall ensure that the PCR states data quality requirements for all data applicable for use in claims. These data shall be verified to be compliant with the established PCR data quality requirements and those for foreground (primary) and background (secondary) data. The PCR shall specify that a data quality assessment be performed on all collected foreground (primary) data and may provide templates to facilitate harmonized primary data collection, assessment, reporting, and verification. <i>Note: Refer to the 'Assessing Data Quality of Background Life Cycle Inventory Datasets' addendum.</i> | This guidance | PCR: • Data quality assessment criteria and/or template | 3 Data source | N/A | N/A |
| ☑ 31 | PCR Committee shall ensure that PCR-designated background (secondary) data sources be specified and verified such that: • Data for electricity, transportation, basic fuels, and heavy equipment operation are the most current versions from common public background data (e.g., for North America, LCI and method compatible with the Federal Elementary Flow List (FEDEFL) from https://www.lcacommons.gov/). • Temporal, geographical, and technological coverage of the secondary data is compatible with the scope of the PCR. • System boundaries are equivalent, and reference flows are adaptable to the product system specified in the PCR. • Sources of secondary data are cited. • Allocation procedures used for secondary data are appropriate for the system under study. | This guidance and 'Assessing Data Quality of Background Life Cycle Inventory Datasets' and the 'Allocating Materials Shared Across Product Systems' addenda | PCR: • Draft PCR with list of background (secondary) data sources and default LCIA method(s) | 2 Procurement | N/A | N/A |
| Allocation | n | | | | How criteria were met | Due |
| ☑ 32 | PCR Committee shall ensure that the PCR specifies which processes are to be subdivided if allocation can be avoided in this manner wherever feasible. The PCR shall also provide guidelines on how the subdivision should be performed. | 14025 Clause 6.7.1c, 6.7.2c 14027 Clause 6.5.3 | PCR • Draft PCR that lists processes and subdivision method | 2 Procurement | N/A | N/A |
| ☑ 33 | PCR Committee shall ensure the PCR specifies that where allocation by physical relationship is applied, the PCR shall specify the relevant underlying physical relationships to be considered and establish or refer to the relevant allocation rules. | 14025 Clause 6.7.1c, 6.7.2c 14027 Clause 6.5.3 | PCR • Draft PCR that includes specification | 1 Transparency | Allocation rules are listed in section 8 of SM Part A. | Complete |
| ☑ 34 | | 14044 Clause 4.3.4 21930 Clause 7.1.7.2.7 | PCR • Draft PCR that includes specification | 1 Transparency | Allocation regarding output of waste per ISO standards is listed in section 8 of SM Part A. | Complete |
| | | | | | | |
| ☑ 35 | PCR Committee shall refer to rules for and prioritize stepwise allocation for industrial processes that produce more than one product or deliver more than one service. For example, the refining of crude oil produces more than one different product, such as liquefied petroleum gas, gasoline, naphtha, diesel, asphalt, and others. PCR Committee shall refer to rules prohibiting system expansion as a method for avoiding allocation for construction products that may involve the production of coproducts; rather, the PCR shall prescribe an ISO-compliant method of allocation, or an allocation procedure if multiple methods are allowed. | 14044 Clause 4.3.4.2 21930 Clause 7.2.5 | PCR • Draft PCR including allocation method and procedure (where applicable) | 2 Procurement | N/A | N/A |

| | | ☑ 36 | PCR Committee shall prescribe ISO-compliant rules for allocation between product systems (across the system boundary) and designate whether Module D may be optionally reported in the EPD for construction products and services. If so, the PCR shall prescribe detailed calculation rules for any quantitative metrics reported therein Note: Refer to the 'Allocating Burdens and Benefits of Materials Shared Across Product Systems'addendum. | 21930 Clause 7.2.6 | PCR: • Draft PCR with allocation rules and calculation rules | 2 Procurement | N/A | N/A |
|------------------------|------|-----------|---|---|--|----------------|--|----------|
| Life cycle im assessme | | ☑ 37 | PCR Committee shall include all minimally required, core indicators for ISO-compliant EPDs; specifically bulleting the indicator with: 1) the LCA characterization methodology, and 2) reference in parenthesis. Additionally, the PCR is encouraged to specify at least one LCIA method that includes characterization factors for calculating category indicator results for each impact category and each geographical region covered by the PCR. | 21930 Clause 9.5 | PCR: • Draft PCR including all items | 1 Transparency | Core indicators are listed in section 9 of SM Part A. | Complete |
| Interpretat | tion | ☑ 38 | PCR Committee shall identify the steps for interpreting the results of the underlying LCA study. | 14044 Clause 4.5 21930 Clause 9 | PCR: • Draft PCR including all items | 1 Transparency | SM Part A section 9.3 includes steps for interpreting the results of a background LCA. | Complete |
| | | ☑ 39 | PCR Committee shall ensure that the PCR communicates requirements (either qualitative or quantitative) and reference the methods and format used to report additional environmental information. | 21930 Clause 8.4 14025 Clause 7.2.3, 7.2.4 | PCR: Detailed specification on requirements and reference methods and format used to report additional environmental information. | 1 Transparency | SM Part A section 10 includes a description of additional environmental information and the TR/EPD template in Appendix C showing placement of such information. | |
| | | ☑ 40 | PCR Committee shall ensure that the PCR lists assumptions and limitations associated with the underlying LCA results. | 14044 Clause 4.5.2.1 | PCR: • Draft PCR including all items | 1 Transparency | SM Part A section 5.2 includes a description of assumptions and limitations associated with TR/EPD results. | Complete |
| | | 41 | PCR Committee shall specify different types of uncertainties to be propagated in the underlying LCA study and is encouraged to ensure that the PCR describes procedures for reporting uncertainty of results. | 14044 Clause 4.4.4.2 14025 6.7.1b | PCR: • Draft PCR including all items | 1 Transparency | SM Part A states that uncertainty shall be addressed in the data quality assessment and may be addressed qualitatively or quantitatively. | Complete |

Part B for: commercial lavatories

October 2, 2024 | Sustainable Minds | Contact Kim Hammer (kim@sustainableminds.com)

EPD use case goal:

1, 2 or 3

EPD use levels are cumulative.
Transparency is the baseline. To create a 'Data source' conformant
PCR, all criteria in all checklists must be documented.

3. PCR Review Panel checklist Version 1.0, May 25, 2022 | ACLCA PCR Open Standard 2022

| Categories | | # | Criteria | ISO reference | Supporting documentation | EPD use | 3 Data source 2 Procurement 1 Transparency | |
|----------------|------|-----|---|---|--|----------------|--|----------|
| | Grou | ınd | rules | | | | How criteria were met | Due |
| | Ø | 1 | The PCR Review Panel shall use this checklist to guide their process of reviewing the PCR. | This guidance | PCR supporting documentation: • Completed checklist | 1 Transparency | Completed checklists saved with the PCR supporting documentation. | Complete |
| | Ø | 2 | PCR Review Panel members shall disclose any conflicts of interest using the conflict of interest form. | 14027 Clause 7.2 14071 | PCR supporting documentation: • Review panel completed conflict of interest forms | 1 Transparency | Conflict of interest forms to be completed by review panel members. | Complete |
| Organizational | Ø | 3 | The PCR Review Panel shall meet with the Program Operator to discuss the PCR and how to perform their review. The PCR Review Panel shall investigate whether the PCR has been developed in accordance with relevant LCA-based claim standards, general program instructions, specifications, and guidelines, and ensure that it supports the creation of credible and consistent claims. The PCR Review Panel shall verify that the EPD template is consistent with the PCR guidelines. The PCR Review Panel shall generate and compile their comments in a review report. By the agreed upon date determined by the Program Operator, the review report shall be sent to the PCR Committee for consideration. | | PCR supporting documentation: • Dated review report | 1 Transparency | Aggregated technical and public comments spreadsheet, including commenter names and committee responses, to be created and made available in the Detailed Review Report. | Complete |
| | Ø | 4 | federal and/or state procurement requirements (e.g., Buy Clean Legislation) that | This guidance and relevant EPD-related federal and/or state procurement requirements | PCR supporting documentation: Reviewers' sign-off and/or list of any deviations from procurement requirements | 2 Procurement | N/A | N/A |
| | Ø | 5 | The PCR Review Panel shall verify conformance the Program Operator and PCR Committee checklists and the appropriate category of EPD use is identified. | This guidance | PCR supporting documentation: • Reviewers' sign-off below and/or list of any deviations from this guidance. All three completed checklists returned to the PO. | 1 Transparency | Section below completed by review panel chair, who confirmed sign-off from all review panel members. | Complete |

Reviewer acceptance for EPD use case (1, 2, or 3) Date | Reviewer names & email

| Date | Revier name & email | Acceptance for EPD use case Level 1 (Y/N) |
|----------|---|---|
| 2-Oct-24 | Thomas Gloria - t.gloria@industrial-ecology.com | Yes |
| 3-Oct-24 | Rifat Karim - rifat.chimique@gmail.com | Yes |
| 2-Oct-24 | Jack Geibig - igeibig@ecoform.com | Yes |



Part B comments worksheet

SM Transparency Report™ Framework

Sustainable Minds, PCR Part B: Product group definition | Commercial Lavatories, 2024. http://www.sustainableminds.com/files/transparency/pgds/Part_B_Product_Group_Definition_Commercial_Lavatories_2024.pdf.

Part B: Product group definition

ame: Commercial lavatories
rs: Tom Gloria, Jack Geibig, Rifat Karim (no comments during public consultation)

| To | pic Page | e Section | Type of comment | Reviewer comment | Reviewer's proposed change/solution | Response | Rationale | Reviewer response to public comment | Response |
|----|----------|---|-----------------------|--|---|-----------|---|--|--|
| 1 | 1 | Product group - Exclusions | Technical | Clear instructions required. | RK: To have a clear instruction, mention that sinks outside of the lavatory are not included in this PCR. Suggested revision as: "Sinks intended for use outside a lavatory (bathroom, toilet room, or bathing facility)" | No change | PCR Committee members have defined lavatory as "a washbowl or basin used in a bathroom, toilet room, or bathing facility" | | Closed |
| 2 | 1 | 2 | tech | PCR states "Any parts/components not integrated and sold with the lavatory or wash fountain shall be excluded from the system boundary." | JG: This is an important criterion, as it determines what is in scope. How is integrated defined here? This seems a little squishy. The word is defined as "with various parts or aspects linked or coordinated." So as written, parts must be both integrated and sold to be included. Are all parts sold with the lav considered intergrated? How can we discern the ones that are and arent? IN short, can this requirment be made simpler by simply saying "all parts sold with the product" or made more specific by defining parts that can be considered integrated? | Accept | Reworded scope to cover lavatory systems and added corresponding definition. | Accept | Closed |
| 3 | 1 | 2 | edit | For sections B6 and B7 it is stated that the rules for lavoratories which include a faucet follow the commercial faucets PCR. | given none of them are sold with the product. While its ok to reference criteria in other guidance, this typically involves guidance that also includes the products within the PCR as a subset (e.g. Referencing 21930 in a PCR for building products). Suggest listing the requirements in this document so the PCR is self contained. | No change | Since the vast majority of lavatories are sold without an integrated faucet, it may be confusing to practitioners to list the lengthy water and energy use assumptions in this Part B as well. When the faucets Part B gets updated, this Part B adopts those assumptions. In addition, Sustainable Minds prefers a modular approach to Part Bs and has used this same method for e.g., the commercial toilets Part B, where B6-B7 refer to the flush valves Part B for integrated flush valves Part B for integrated flush valve products. | Accept | Closed |
| 4 | 1 | 3 | edit | PCR states " An existing non-expired PCR for commercial lavatories was not found at the time of publication of this Part B " | JG: Awkward sentence. Rewrite as "At the time of publication of this Part B, an active PCR for commercial lavatories was not identified" | Accept | Reworded for clarification as suggested | Accept | Closed |
| 5 | 3 | System boundary | Technical / Editorial | Suggestion for improvement | RK: I suggest re-wording part of the second paragraph. The significance of capital goods and infrastructure to the overall impacts of the products is quantified differently and they vary based on primary and secondary datasets and the dataset providers. To reduce possible artificial variation in EPD results across the product group, capital goods and system infrastructure flows shall be excluded from the system boundary by default, with justification required for alternative assumptions.* | Accept | Reworded second paragraph to improve clarity, implementing phrases suggested. | Accept | Closed |
| 6 | 3 | Additional rules for comparabili ty (A2, A5) | Technical | There are a number of truck transportation option based on the dataset/software is used, since this PCR is based on North America we should specify the truck type following EPA classification. | RK: Type of truck /class needs to be specified. | Accept | Added specification of class 8 truck for both A2 and A4. Reference: https://afdc.energy.gov/data/10380 | Accept | Closed |
| 7 | 3 | | tech | PCR States "EPDs that use secondary data for any unit process that contributes 5% or more to any disclosed environmental impact category shall disclose the data source (database name and version, software type and version implemented, dataset name, dataset geography, and dataset allocation method)." | JG: I noticed this additional reporting requirement that was not in previous PCRs was added, DO you mean LCA modeling software type? Pls clarify to what type of software this is referring. Also, it should be made clear in the language that the criterion applies to the LCI being used, and not the actual unit process data being reported by the manufacturer. The value of 5% is really low in this instance and will basically require the reporting of details for individual LCI data sets in the EPD itself, rather than just in the LCA background report. | No change | The committee confirmed that 5% is appropriate. Left as-is. | Since the information required to be reported is relevant to the data source, it remains unclear to what "software type and version" is referring? LCI data are often available outside the structure of a software tool (eg. ecoinvent data can be purchased separately from multiple LCA tools). This specific requirement should be explained or deleted. | Changed to "LCI database name and version" and "LCA modeling software type and version implemented" This specific requirement was added in response to a previous PCR Committee comment that there's a chance the same data set may provide different results depending on software used. JG: Accepted |
| 8 | 3 | Additional rules for comparabili ty (A4) | Editorial | Suggestion for improvement | RK: The last sentence is a little bit confusing. Suggestion: "1600km (994 miles) should be the total distance inclusive of several transport legs (if any)." | Accept | Reworded second sentence to clarify total of 1600km and that it includes multiple transport legs if needed. | Accept | Closed |
| 9 | 4 | 2 | edit | In the A1 section it is stated that "the electricity grid profile of the data set should be adapted to the source country or region, if known and possible with the selected data set." | JG & TG: Replace should with shall. | Accept | Shall statement aligns with the intent of this scenario | Accept | Closed |

| 10 | 4 | Additional rules for comparabili ty (A4) | Editorial | Additional character | RK: Please remove the additional "(" in front of 994 | Accept | Reworded second sentence to clarify total of 1600km and that it includes multiple transport legs if needed. | Accept | Closed |
|----|---|--|-----------|--|--|------------------|---|--|---|
| 11 | 4 | 2 | edit | PCR States "In cases when the EPD owner purchases manufacturing process activity at the upstream supplier shall be counted in the extraction and upstream production stage, separate and in addition to the upstream raw material extraction." | JG: This language is unclear and confusing. By "extraction and upstream production" do you mean the raw material extraction and processessing stage? This should be clarified. The phrase "separate and in addition to" is really unclear. Separate from what, the extraction stage (which seems to contradict the first part of the sentence requiring its inclusion there?) In addition, to what? Is this suggesting that the values fro upstream parts manufactruing be reported separate and in addition to other raw material extraction? I dont get this at all. | Accept | Reworded to: "In cases when the EPD owner purchases manufactured components, the manufacturing process activity at the upstream supplier shall be counted in this stage, in addition to the upstream raw material extraction." | Accept | Closed |
| 12 | 4 | Additional rules for comparabili ty (A5) | Technical | The sentence is unclear | RK. Add related to installation at the end of the sentence. "The installation stage shall include, as applicable, any ancillary materials, electricity and/or water consumption (e.g., from tools or initial product testing by customer prior to lirst use), and disposal of product packaging waste and other waste materials directly related to the installation phase." | Accept | Added "directly related to installation of the product." for clarity. | Accept | Closed |
| 13 | 4 | 2 | edit | In the B3 section, a number of different scenarios are broken out for repairs for the lavoratories. However, it should also make clear that when there are no drains or faucets sold with the product then repairs are not necessary (zero). In other sections, like B6 / B7 this is broken out. It needs to be broken out here | JG: Include a clear statement that repairs are not applicable when a product is sold without integrated components like a drain or a faucet. | Accept | Added intro to state that repairs are not expected for lavatories without a faucet or drain. | Accept | Closed |
| 14 | 4 | Additional rules for comparabili ty (Reference service life) | Technical | Calrity in instructions required. | RK: Please provide a conversion caltulation/rule between RSL, ESL and reference flow (described in B3, can be copied/moved/referenced). | No change | The ESL and RSL section is intended to specify the ESL and RSL, which are relevant for the assumptions laid out in B1-B7. The conversion calculation/rule in B4 is only relevant for replacements. | Accept | Closed |
| 15 | | Additional rules for comparabili ty (Maintenan ce B2) | Technical | Since there is a variation in how the lavatories are sold in the North American market (with or without faucets), the claning activity will change. | RK: Provide clear guidance on cleaning scenarios. | Accept | Referred to faucets Part B for additional cleaning for products with integrated faucets and added table for quicker reference. | Accept | Closed |
| 16 | 4 | Additional rules for comparabili ty (Repair B3) | Technical | This section should discuss only the repair and not the replacement. | RK: Move the replacement discussion to the next section (B4) | Slight rewording | A faucet is a repairable component of lavatory systems (definition added for clarity) and should be replaced every 10 years to align with the faucet PCR. Slightly edited wording for clarity but left mostly as-is. | Accept | Closed |
| 17 | 4 | Additional rules for comparabili ty / Default Life Cycle Stage Scenarios (Replaceme nt B4) | Editorial | | JG & TG: Use this example to clearly and unambiguously demonstrate that the number of replacements of 2.75, should be rounded to the nearest tenth, as in 2.8. | Accept | Clarified to the nearest hundredth. | Accept | Closed |
| 18 | 4 | Additional rules for comparabili ty (B6 & B7) | Technical | To avoid going back and fort between PCRs can we put the reference text here? | RK: Add the reference text here from the commercial/public metered and manual lavatory faucets here. Or at the very least external link / the section numbers should be added. | No change | Since the vast majority of lavatories are sold without an integrated faucet, it may be confusing to practitioners to list he lengthy water and energy use assumptions in this Part B as well. When the faucets Part B gets updated, this Part B adopts those assumptions. In addition, Sustainable Minds prefers a modular approach to Part Bs and has used this same method for e.g., the commercial toilets Part B, where B6-B7 refer to the flush valves Part B for integrated flush valve products. | Agree. Missing the closing quotation mark. Please fix. | Added closed quotation mark after 'Default life cycle stage scenario(s)' RK: Accept |
| 19 | 5 | Additional rules for comparabili ty (Waste processing C3) | Editorial | Being specific for EoL guidance | RK: To be more clear and specific I suggest changing the last senctence in C3 a little bit to: "take-back program require evidence such as documentation of the program and documented number or share of units sold that participate in take-back program." | Accept | Added clarification to define program as take-back program | Accept | Closed |

| 20 | 5 | | Location for end of life disposition is not fully described. | TG: Provide guidance on default location of NA for end of life disposition if unknown and report if known. | | Added North America for default landfill location | Accept | Closed |
|----|---|---|---|--|----------------------------------|--|--------|--------|
| 21 | 5 | Additional LCA calculation rules | 2017 is the latest updated version for ISO 21930 | RK: Please change the ISO name to "ISO 21930:2017", also to be consistent with the system boundary description. | Accept | Added version to standard | Accept | Closed |
| 22 | 5 | Part B developmen t information (Part B review pane;) | Change Rifat's affiliation | RK: Please change my company name as "Independent Consultant" | Accept | Updated affiliation | Accept | Closed |
| 23 | * | Technical | Clarity with renewable electricity | RK: While I am aware that the REC addendum is currently ongoing through ACLCA and assuming it will be reflected in part A either with revisions or addendum, we need to reference that in this document. | Accepted for inclusion in Part A | RECs are currently not addressed, but they will be in the next SM Part A update, intended to align with the updated ACLCA PCR Open Standard REC addendum once published. | | Closed |
| 24 | 5 | information (Update justification) | This is a new PCR (NOT an update), the justification statement should reflect that. | RK: Suggest changing the word from "updated" to "created". | Accept | Justification for update not needed | Accept | Closed |
| 25 | 5 | Part B developmen t information (Part B review pane;) | Update referencing ISO 21930 to ISO 21930:2017 | RK: The last addendum of ISO 21930 is 2017, either we add the 2017 at the end or drop it to be consistent throughout the document. [also see comment #12] | Accept | Added version to standard | Accept | Closed |