

| PP4   | <p>Patient's phenotype or family history is highly specific for a disease with a single genetic etiology.</p> <ul style="list-style-type: none"> <li>PP4 applicability and strength is determined by the total points accumulated by a single affected individual according to the table below and the following total point ranges: <ul style="list-style-type: none"> <li>&lt;1 point: PP4 not met</li> <li>1-&lt;2 points: PP4</li> <li>≥2-&lt;4 points: PP4_Moderate</li> <li>≥4 points: PP4_Strong<sup>1</sup></li> </ul> </li> </ul> <p><sup>1</sup>CNV (Copy number variation) testing is required to consider PP4_Strong in order to certify that the variant in question is causative for the phenotype, and not one CNV event corrected by gene therapy and not identified previously.</p> <table border="1"> <thead> <tr> <th>Evidence Description</th><th>Points</th></tr> </thead> <tbody> <tr> <td>Diagnostic criteria for SCID/Leaky SCID/Omenn syndrome met<sup>2</sup></td><td>0.5</td></tr> <tr> <td>SCID gene panel or exome/genome sequencing conducted (only applicable if genetic testing did not provide an alternative genetic explanation for SCID/Leaky SCID/Omenn syndrome phenotype)</td><td>0.5</td></tr> <tr> <td>Family history of SCID (only applicable if SCID gene panel or exome/genome sequencing was conducted on proband and did not provide an alternative genetic explanation for phenotype)</td><td>0.5</td></tr> <tr> <td>Decreased presence of TCRVα7.2 in CD3+ T lymphocytes and/or mucosa-associated invariant T-cells demonstrated by flow cytometry</td><td>0.5</td></tr> <tr> <td>SCID phenotype corrected by RAG1 gene therapy <b>WITHOUT</b> CNV testing performed</td><td>1</td></tr> <tr> <td>SCID phenotype corrected by RAG1 gene therapy <b>WITH</b> CNV testing performed</td><td>4</td></tr> <tr> <td>T-B-NK+ lymphocyte subset profile* (See notes)</td><td>0.5</td></tr> </tbody> </table> <p><sup>2</sup>The diagnostic criteria should follow the PIDTC 2022 specification, summarized <a href="#">here</a>. *Notes: 1) If NK cells are not noted or are present, criteria may still be applied if SCID gene panel or exome/genome sequencing has ruled out alternative causes; 2) If maternal T cells are present, the T lymphocyte profile is still considered to be T- (autologous T cells are absent).</p> | Evidence Description | Points | Diagnostic criteria for SCID/Leaky SCID/Omenn syndrome met <sup>2</sup> | 0.5 | SCID gene panel or exome/genome sequencing conducted (only applicable if genetic testing did not provide an alternative genetic explanation for SCID/Leaky SCID/Omenn syndrome phenotype) | 0.5 | Family history of SCID (only applicable if SCID gene panel or exome/genome sequencing was conducted on proband and did not provide an alternative genetic explanation for phenotype) | 0.5 | Decreased presence of TCRVα7.2 in CD3+ T lymphocytes and/or mucosa-associated invariant T-cells demonstrated by flow cytometry | 0.5 | SCID phenotype corrected by RAG1 gene therapy <b>WITHOUT</b> CNV testing performed | 1 | SCID phenotype corrected by RAG1 gene therapy <b>WITH</b> CNV testing performed | 4 | T-B-NK+ lymphocyte subset profile* (See notes) | 0.5 |
|---|---|----------------------|--------|---|-----|---|-----|--|-----|--|-----|--|---|---|---|--|-----|
| Evidence Description  | Points  |                      |        |   |     |   |     |  |     |  |     |  |   |   |   |  |     |
| Diagnostic criteria for SCID/Leaky SCID/Omenn syndrome met <sup>2</sup>   | 0.5   |                      |        |   |     |   |     |  |     |  |     |  |   |   |   |  |     |
| SCID gene panel or exome/genome sequencing conducted (only applicable if genetic testing did not provide an alternative genetic explanation for SCID/Leaky SCID/Omenn syndrome phenotype) | 0.5   |                      |        |   |     |   |     |  |     |  |     |  |   |   |   |  |     |
| Family history of SCID (only applicable if SCID gene panel or exome/genome sequencing was conducted on proband and did not provide an alternative genetic explanation for phenotype)      | 0.5   |                      |        |   |     |   |     |  |     |  |     |  |   |   |   |  |     |
| Decreased presence of TCRVα7.2 in CD3+ T lymphocytes and/or mucosa-associated invariant T-cells demonstrated by flow cytometry  | 0.5   |                      |        |   |     |   |     |  |     |  |     |  |   |   |   |  |     |
| SCID phenotype corrected by RAG1 gene therapy <b>WITHOUT</b> CNV testing performed  | 1   |                      |        |   |     |   |     |  |     |  |     |  |   |   |   |  |     |
| SCID phenotype corrected by RAG1 gene therapy <b>WITH</b> CNV testing performed   | 4   |                      |        |   |     |   |     |  |     |  |     |  |   |   |   |  |     |
| T-B-NK+ lymphocyte subset profile* (See notes)  | 0.5   |                      |        |   |     |   |     |  |     |  |     |  |   |   |   |  |     |