Conditions of Use.

1.1 Prior to use of the data and/or bio-specimens, the Researcher will have obtained approval of the project by an appropriate HREC and research governance/SSA.

1.2 The Researcher must comply with all applicable privacy legislations in relation to the use, disclosure and storage of the data and/or biospecimens.

1.3 The requested data and/or biospecimens are consistent with the research protocol approved by the HREC.

1.4 The data will be stored in a secure place (e.g., password lock on electronic data), the user name and password will be secure and not shared.

1.5 The data and/or biospecimens will be analysed, stored, retained and disposed of in accordance with the research protocol approved by the HREC. In addition, any other governing bodies and legislative requirements.

1.6 Keep the non-identifiable patient ID with the record so that the subject can be re-identified, if required for ethical reasons.

1.7 The data and/or biospecimens released may only be used by researchers under the supervision of the approved Principal Investigator for the project, for the term of the project. The Researcher must not use the data for any other purpose.

1.8 Never publish or communicate data where the sample size and data would allow unintended identification.

1.9 Applicants must ensure that their expression of interest and application to access materials are complete.

2.0 Researchers must satisfy ABTB members that their research projects are ethically and scientifically justified.

2.1 Any materials released to the Researcher may by subject to a Materials Transfer Agreement (MTA) between the Research and ATBA member supplying materials.

2.2 The Researcher must ensure any remaining materials after completion of the project are destroyed or returned as agreed by the relevant ABTB member/s.

2.3 All costs of retrieval, processing and transportation of materials and any other costs associated with the study will be met by the Researcher unless other arrangement have been made with ABTB member/s.

2.4 The Researcher must acknowledge the contribution of ABTB and its funders from which the materials originated in all publications resulting from the use of materials.

2.5 Co-authorship on resultant publications due to any contributions to a specific project is directly between the individual biobank member/s and the Principal Investigator. It is expected that the International Committee of Medical Journal Editors (ICMJE) are followed.

2.6 ABTB can offer no guarantee that materials are devoid of infectious agents.
2.7 ABTB tumour samples are categorised by the diagnosis given by our Histopathologists.

2.8 It is the responsibility of the Researcher to ensure that all staff are appropriately informed and trained in relation to the dangers and procedures in the handling of human materials and that they are trained in, and adhere to, local conditions and Occupational Health and Safety regulations.

2.9 No responsibility will be taken by ABTB for an injury, damages or loss that may occur as a direct or indirect result of the use of materials provided.

3.0 The Researcher agrees to assume all responsibility and risks for the receipt, storage, handling and use of all materials and/or data obtained from ABTB.

3.1 The Researcher agrees to submit annual reports to the ABTB Coordinator and report to the relevant member biobank/s as per their protocol.

Please note these terms and conditions are subject to change.