Table S1. **Default** abagen **pipeline options** | The default settings for the 17 processing steps considered when processing the AHBA data with abagen. An entry of "—" indicates that this is a required, user-supplied parameter. A blank entry indicates that the processing step is not implemented by default. Refer to Table **??** and *Methods: Gene expression pipelines* for further details.

Option	Default
Volumetric or surface atlas	_
Individualized or group atlas	_
Use non-linear MNI coordinates	True
Mirror samples across L/R hemisphere	
Update probe-to-gene annotations	True
Intensity-based filtering threshold	50%
Inter-areal similarity threshold	
Probe selection method	differential stability
Donor-specific probe selection	aggregate
Missing data method	
Sample-to-region matching tolerance	2mm
Sample normalization method	scaled robust sigmoid
Gene normalization method	scaled robust sigmoid
Normalize only matched samples	True
Normalizing discrete structures	False
Sample-to-region combination method	donors
Sample-to-region combination metric	mean