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MIDUS DNA Methylation Age Data

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Abstract

MIDUS is releasing Methylation Age scores data derived from DNA. See associated documentation for details.

Title

MIDUS DNA Methylation Age Data

Related Materials

Other

- Readme
- Documentation

M2MR1_GEN_DNAmAge_N2118_20221027

Title	M2MR1_GEN_DNAmAge_N2118_20221027		
File Name	M2MR1_GEN_DNAmAge_N2118_20221027.sav		
Case Quantity	2118		
Variable Count	19		

M2MRID - M2 AND MR Public IDs combined in a single variable

Type	Numeric (Integer)
Display Values	2
Stats	1

Valid	Invalid	Minimum	Maximum	Mean	StdDev
2118	0	10002	39977	23028.720	10360.625

SAMPLMAJ - Major sample identification

Туре	Code
Display Values	1
Stats	1

			Frequency	% of total	% of valid
Valid	1	CORE MAIN RDD	640	30.22%	30.22%
	2	CORE SIBLING	6	0.28%	0.28%
	3	CORE TWIN	388	18.32%	18.32%
	4	CORE CITY OVERSAMPLE	20	0.94%	0.94%
	13	CORE MILWAUKEE	201	9.49%	9.49%
	20	REFRESHER MAIN	746	35.22%	35.22%
	21	REFRESHER MILWAUKEE	117	5.52%	5.52%
		Total	2,118	100%	100%

Valid	Invalid	Minimum	Maximum
2118	0	1	21

M2MRCASE - Which sample is R in? 1=M2, 2=MR1 Markon Market Sample is R in? 1=M2, 2=MR1 Market Marke

Туре	Code
Display Values	1
Stats	1

			Frequency	% of total	% of valid
Valid	1	M2	1,255	59.25%	59.25%
	2	MR1	863	40.75%	40.75%
	3	M3 NEWTWIN		0%	0%
		Total	2,118	100%	100%

Valid	Invalid	Minimum	Maximum
2118	0	1	2

SENCONSENT - Did R consent to genetics 1=Yes, 2=No

Туре	Code
Display Values	1
Stats	1

			Frequency	% of total	% of valid
Valid	1	YES	2,009	94.85%	94.85%
	2	NO	109	5.15%	5.15%
		Total	2,118	100%	100%
Missing	9	INAPP-M3NEWTWIN		0%	
		Total	0	0%	

Valid	Invalid	Minimum	Maximum
2118	0	1	2

SECOND SECOND S

Туре	Code
Display Values	1
Stats	1

			Frequency	% of total	% of valid
Valid	0	NONE	229	10.81%	10.81%
	1	BLOOD	1,418	66.95%	66.95%
	2	SALIVA	471	22.24%	22.24%
		Total	2,118	100%	100%
Missing	9	INAPP-M3NEWTWIN		0%	
		Total	0	0%	

Valid Invalid	Minimum	Maximum	
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2118	0	0	2

SBRA6DMAVAIL - Is Methylation score data available?

Туре	Code
ForwardSkip	IF <u>BRA6DMAVAIL</u> = 2, SKIP <u>BRA6DMQCCORR</u> TO <u>BRA6DMAGEDUNEDINPACE</u>
Display Values	1
Stats	1

			Frequency	% of total	% of valid
Valid	1	YES	1,310	61.85%	61.85%
	2	NO	808	38.15%	38.15%
		Total	2,118	100%	100%
Missing	9	INAPP-M3NEWTWIN		0%	
		Total	0	0%	

Valid	Invalid	Minimum	Maximum
2118	0	1	2

SBRA6DMQCCORR - Correlation between the sample DNAm profile and the

gold standard reference profile

Туре	Numeric (Double)
Numeric Details	Decimals: 4
BackwardSkip	(BRA6DMAVAIL = 2)
Display Values	2
Stats	2

			Frequency	% of total	% of valid
Missing	9.0000	INAPP	808	38.15%	
		Total	808	38.15%	

Valid	Invalid	Minimum	Maximum	Mean	StdDev
1310	808	0.9306	0.9744	0.9698843	0.0027901

SBRA6DMQCDIF - Difference the sample DNAm profile from the gold

standard reference profile

Туре	Numeric (Double)
Numeric Details	Decimals: 4
BackwardSkip	(BRA6DMAVAIL = 2)
Display Values	2
Stats	2

			Frequency	% of total	% of valid
Missing	9.0000	INAPP	808	38.15%	

Total				808	38.15%	
Valid	Invalid	Minimum	Maximum	Mear	1	StdDev
1310	808	0.051	0.0795	0.054	6251	0.0022431

SRA6DMQCFEMALE - Participant sex as inferred from X chromosome methylation abundance

Туре	Numeric (Integer)
BackwardSkip	(BRA6DMAVAIL = 2)
Display Values	1
Stats	1

			Frequency	% of total	% of valid
Missing			1	0.05%	
	9	INAPP	808	38.15%	
		Total	809	38.2%	

Valid	Invalid	Minimum	Maximum	Mean	StdDev
1309	809	0	1	0.5576776	0.4968519

SECOND SECOND S

Туре	Numeric (Double)
Numeric Details	Decimals: 4
BackwardSkip	(BRA6DMAVAIL = 2)
Display Values	2
Stats	2

			Frequency	% of total	% of valid
Missing	9.0000	INAPP	808	38.15%	
		Total	808	38.15%	

Valid	Invalid	Minimum	Maximum	Mean	StdDev
1310	808	0.3065	0.5435	0.4092324	0.0755935

SBRA6DMARRAYID - Individual microarray used to assay samples

Туре	Text
BackwardSkip	(BRA6DMAVAIL = 2)
Display Values	3
Stats	3

Valid	Invalid
2118	0

SBRA6DMPLATE - Individual 96-well plate storing samples for assay

Туре	Numeric (Integer)
BackwardSkip	(BRA6DMAVAIL = 2)
Display Values	2
Stats	1

			Frequency	% of total	% of valid
Missing	99	INAPP	808	38.15%	
		Total	808	38.15%	

Valid	Invalid	Minimum	Maximum	Mean	StdDev
1310	808	1	14	7.4923664	4.0205187

SRA6DMWELL - Individual well within the 96-well plate

Туре	Text
BackwardSkip	(BRA6DMAVAIL = 2)
Display Values	3
Stats	3

Valid	Invalid
2118	0

SBRA6DMAGEHORVATH - DNA methylation age - Horvath

Туре	Numeric (Double)
Numeric Details	Decimals: 4
BackwardSkip	(BRA6DMAVAIL = 2)
Display Values	2
Stats	2

			Frequency	% of total	% of valid
Missing	999.0000	INAPP	808	38.15%	
		Total	808	38.15%	

Valid	Invalid	Minimum	Maximum	Mean	StdDev
1310	808	25.3443	107.2746	55.460805	11.118578

SBRA6DMAGEHORVATH2 - DNA methylation age - Horvath2 (Skin) €

Туре	Numeric (Double)
Numeric Details	Decimals: 4
BackwardSkip	(BRA6DMAVAIL = 2)
Display Values	2
Stats	2

		Frequency	% of total	% of valid
Missing	999.0000 INAPP	808	38.15%	

	Total			808	38.15%	
Valid	Invalid	Minimum	Maximum	Mean	l	StdDev
1310	808	18.8184	98.3801	51.90	6437	12.444525

₱ BRA6DMAGEHA	NNUM - DNA methylation age - Hannum
Туре	Numeric (Double)
Numeric Details	Decimals: 4
BackwardSkip	(BRA6DMAVAIL = 2)
Display Values	2
Stats	2

			Frequency	% of total	% of valid
Missing	999.0000	INAPP	808	38.15%	
		Total	808	38.15%	

Valid	Invalid	Minimum	Maximum	Mean	StdDev
1310	808	14.0105	110.4416	42.441519	11.456398

SRA6DMAGEPHENOAGE - DNA methylation age - PhenoAge

Туре	Numeric (Double)
Numeric Details	Decimals: 4
BackwardSkip	(BRA6DMAVAIL = 2)
Display Values	2
Stats	2

			Frequency	% of total	% of valid
Missing	999.0000	INAPP	808	38.15%	
		Total	808	38.15%	

Valid	Invalid	Minimum	Maximum	Mean	StdDev
1310	808	10.2689	85.0056	43.636760	12.994821

SBRA6DMAGEGRIMAGE - DNA methylation age - GrimAge

Туре	Numeric (Double)
Numeric Details	Decimals: 4
BackwardSkip	(BRA6DMAVAIL = 2)
Display Values	2
Stats	2

			Frequency	% of total	% of valid
Missing	999.0000	INAPP	808	38.15%	
		Total	808	38.15%	

Valid	Invalid	Minimum	Maximum	Mean	StdDev
1310	808	22.497	104.5895	52.649525	11.125769

BRA6DMAGEDUNEDINPACE - DNA methylation age - DunedinPACE Type Numeric (Double) Numeric Details Decimals: 4 BackwardSkip (BRA6DMAVAIL = 2) Display Values 2

		Total			808	38.15%	
Missing	9.0000	INAPP			808	38.15%	
					Frequency	% of total	% of valid

2

Stats

Valid	Invalid	Minimum	Maximum	Mean	StdDev
1310	808	0.5336	1.4541	0.9909680	0.1374513