

Fig.S1. The average number of sites corresponding to different proportions of missing values on all cancers

Supplementary Figure 2

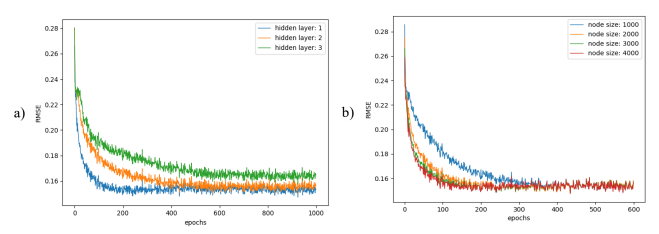


Fig.S2. Selection process of hidden layer and node size. The x coordinate is the training epoch, and the y coordinate is the training loss. (a) Shows that when the hidden layer is 1, the convergence speed is faster. (b) Shows that the RMSE of the node size 3000 and 4000 are close, the value of the node size is determined to be 3000 to reduce some calculations.

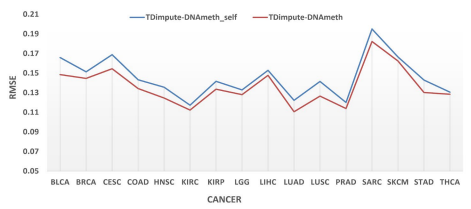


Fig.S3. Comparing the RMSE of TDimpute-DNAmeth_self and TDimpute-DNAmeth methods on 16 cancers. The results were averaged over four missing rates. Table S3 details the RMSE value at the four missing rates.

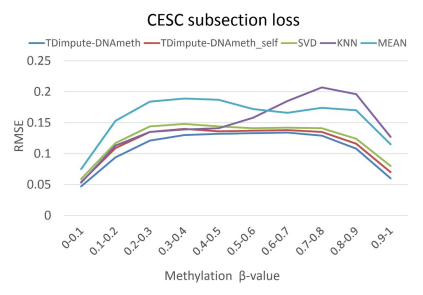


Fig.S4. Subsection RMSE of five methods on CESC cancer datasets. There is no obvious relationship between the β -value of methylation and the RMSE.

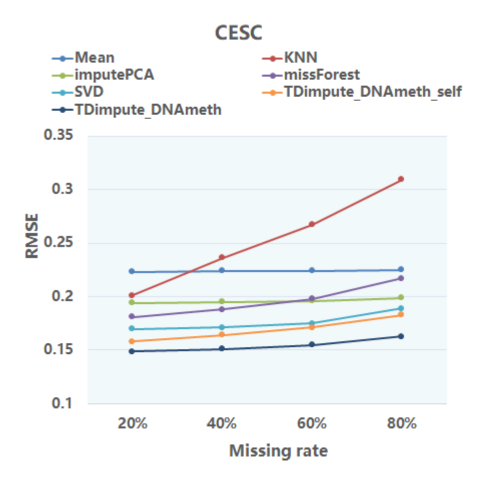


Fig.S5. The comparison of seven methods by RMSE on the CESC dataset. The missForest is not as effective as the SVD method, and the time taken by this method is unacceptable. Therefore, the missForest is not in the manuscript.

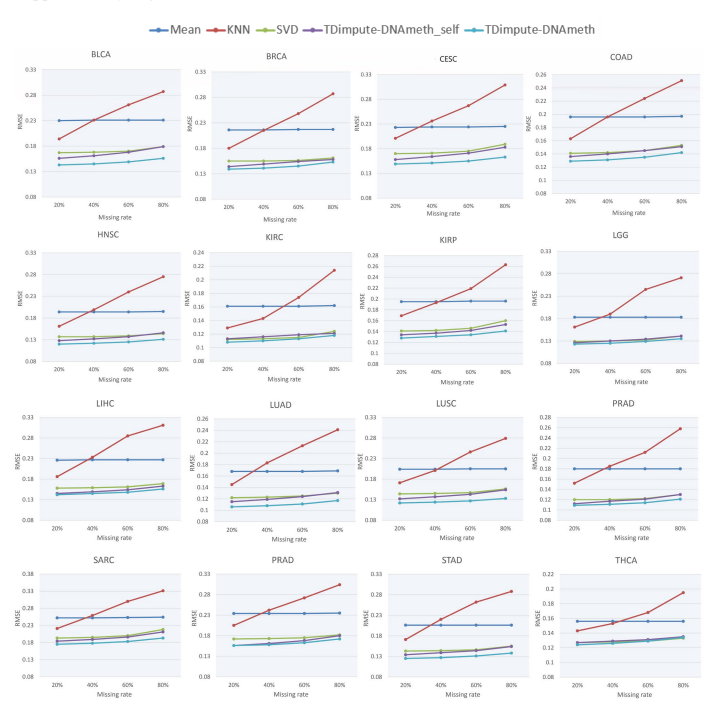


Fig.S6. RMSE of five methods on 16 imputed cancer datasets with different missing rates. The results were averaged over 5 random replicas.

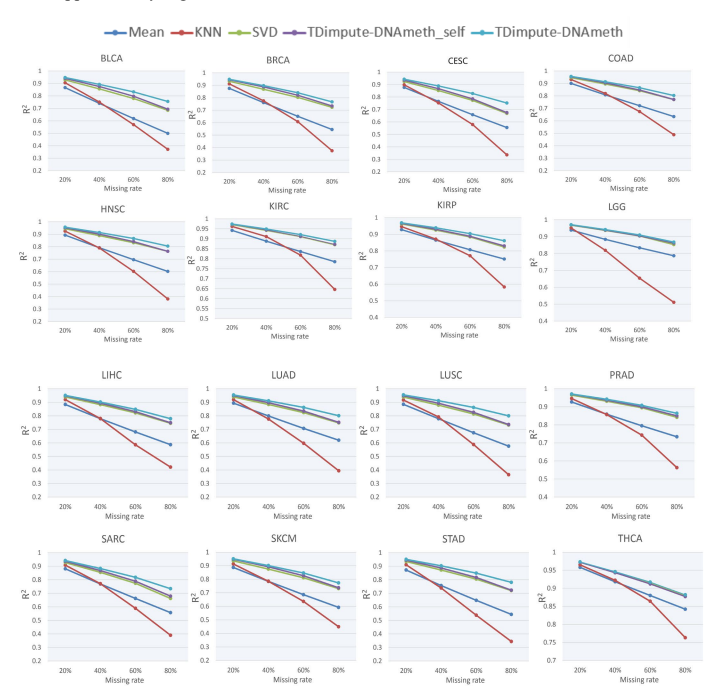


Fig.S7. R² of five methods on 16 imputed cancer datasets with different missing rates. The results were averaged over 5 random replicas.

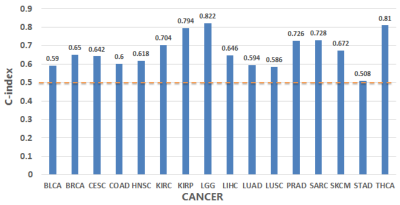


Fig.S8A. The C-index of the survival prediction of 16 cancers on the TCGA dataset without simulation missing values.

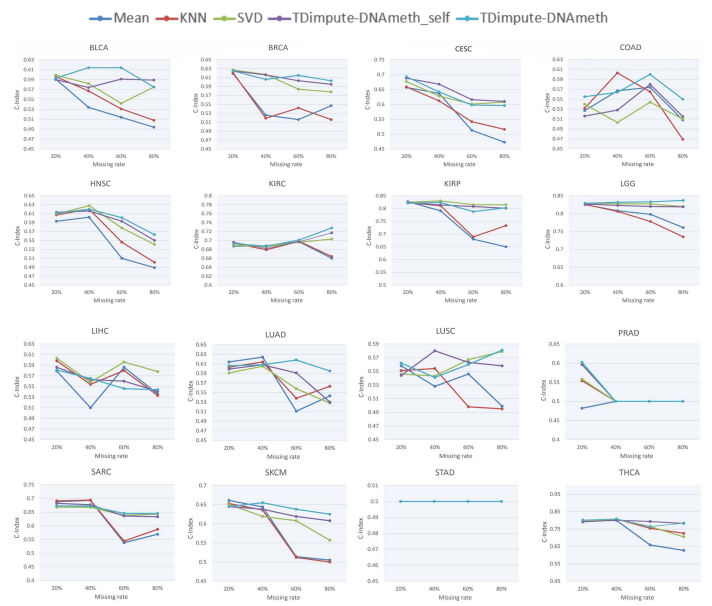


Fig.S8B. C-index of five methods on 16 imputed cancer datasets with different missing rates. The results were averaged over 5 random replicas. If the predicted value of all methods is less than 0.5, use 0.5 instead.

Supplementary Table

	. The RWISE C	BLC		6 cancer datase		BRCA		
Miss-	TDi		.A TDi			BRCA	TDin	
ing	pute-DNA		pute-DN		TDimpute-DN	Ameth_self	pute-DN	
rate	cv(5fold)	test	cv(5fold)	test	cv(5fold)	test	cv(5fold)	test
20%	0.1584± 0.0052	0.1581	0.1481 ±0.0029	0.1488	0.1443 ± 0.0029	0.1425	0.1404± 0.0029	0.1392
40%	0.1647± 0.0051	0.1647	0.1518 ±0.0029	0.1524	0.1523 ± 0.0029	0.1511	0.1447 ± 0.0027	0.1436
60%	0.1735± 0.0051	0.1733	0.1572 ±0.0030	0.1574	0.1627 ± 0.0030	0.1619	0.1502 ± 0.0026	0.1490
80%	0.1889± 0.0054	0.1881	0.1677± 0.0032	0.1676	0.1768 ± 0.0032	0.1758	0.1591 ± 0.0027	0.1578
Miss-		CES	C			COAD		i
	TDi	m-	TDi	m-	TDimmita DN	A ath - a alf	TDin	n-
ing	pute-DNA	meth_self	pute-DN	Ameth	TDimpute-DN	Amem_sen	pute-DNA	Ameth
rate	cv(5fold)	test	cv(5fold)	test	cv(5fold)	test	cv(5fold)	test
20%	0.1653± 0.0047	0.1646	0.1514± 0.0040	0.1516	0.1387 ± 0.0053	0.1374	0.1333± 0.0047	0.1324
40%	0.1725± 0.0047	0.1710	0.1559± 0.0039	0.1558	0.1451 ± 0.0051	0.1449	0.1373 <u>±</u> 0.0048	0.1366
60%	0.1839± 0.0052	0.1796	0.1619± 0.0040	0.1616	0.1542± 0.0051	0.1553	0.1421± 0.0048	0.1416
80%	0.2059± 0.0058	0.1986	0.1737± 0.0043	0.1736	0.1701± 0.0051	0.1706	0.1514± 0.0048	0.1510
Miss-		HNS	SC			KIRC		i
ing	TDi	m-	TDi	m-	TDimmita DN	A ath - a alf	TDim-	
•	pute-DNA	meth_self	pute-DN	Ameth	TDimpute-DN	Amem_sen	pute-DNAmetl	
rate	cv(5fold)	test	cv(5fold)	test	cv(5fold)	test	cv(5fold)	test
20%	0.1312± 0.0036	0.1277	0.1236± 0.0031	0.1208	0.1122± 0.0047	0.1100	0.1073± 0.0037	0.1060
40%	0.1374± 0.0036	0.1353	0.1271± 0.0032	0.1242	0.1168± 0.0048	0.1178	0.1104 <u>+</u> 0.0037	0.1092
60%	0.1468± 0.0037	0.1460	0.1322± 0.0032	0.1294	0.1246± 0.0050	0.1302	0.1141± 0.0036	0.1128
80%	0.1634± 0.0039	0.1608	0.1416± 0.0035	0.1380	0.1363± 0.0050	0.1412	0.1209± 0.0040	0.1190
Miss-		KIR	P			LGG		<u></u>
ing	TDi pute-DNA		TDi pute-DN		TDimpute-DN	Ameth_self	TDin pute-DN	
	cv(5fold)	test	cv(5fold)	test	cv(5fold)	test	cv(5fold)	test
rate	0.(01010)		0 1010 1		0.1284±	0.1263	0.1275± 0.0024	0.1260
rate 20%	0.1405± 0.0071	0.1369	0.1319± 0.0052	0.1314	0.0026		0.0024	
	0.1405±	0.1369	1	0.1314	0.0026 0.1339± 0.0026	0.1320	0.1315±	0.1296
20%	0.1405± 0.0071 0.1460± 0.0076 0.1539±		0.0052 0.1355± 0.0053 0.1403±		0.1339± 0.0026 0.1421±	0.1320	0.1315± 0.0023 0.1359±	0.1296 0.1346
20%	0.1405± 0.0071 0.1460± 0.0076 0.1539± 0.0081 0.1705±	0.1424	0.0052 0.1355± 0.0053 0.1403± 0.0055 0.1504±	0.1346	0.1339± 0.0026 0.1421± 0.0028 0.1606±		0.1315± 0.0023 0.1359± 0.0022 0.1435±	
20% 40% 60%	0.1405± 0.0071 0.1460± 0.0076 0.1539± 0.0081	0.1424 0.1510	0.0052 0.1355± 0.0053 0.1403± 0.0055 0.1504± 0.0061	0.1346 0.1400	0.1339± 0.0026 0.1421± 0.0028	0.1398	0.1315± 0.0023 0.1359± 0.0022	0.1346

rate	pute-DNA	meth_self	pute-DN	Ameth			pute-DN	Ameth
	cv(5fold)	test	cv(5fold)	test	cv(5fold)	test	cv(5fold)	test
20%	0.1533	0.1517	0.1485	0.1472	0.1179	0.1189	0.1103	0.1122
40%	0.1598	0.1581	0.1526	0.1510	0.1295	0.1298	0.1138	0.1158
60%	0.1679	0.1666	0.1579	0.1564	0.1440	0.1438	0.1184	0.1206
80%	0.1814	0.1797	0.1674	0.1656	0.1573	0.1568	0.1270	0.1286
Miss-		LUS	C			PRAD		•
ing	TDi pute-DNA		TDi		TDimpute-DNA	Ameth_self	TDir	
rate	cv(5fold)	test	pute-DN cv(5fold)	test	cv(5fold)	test	pute-DN/ cv(5fold)	test
	0.1385±	test	0.1268±	iesi	0.1168±	test	0.1153±	test
20%	0.0056	0.1370	0.0050	0.1264	0.0028	0.1124	0.0025	0.1118
40%	0.1454± 0.0057	0.1441	0.1303± 0.0048	0.1300	0.1246± 0.0028	0.1199	0.1197± 0.0024	0.1162
60%	0.1570± 0.0059	0.1555	0.1355± 0.0050	0.1352	 0.1374± 0.0030	0.1320	0.1251± 0.0025	0.1214
80%	0.1733± 0.0063	0.1714	0.1456± 0.0054	0.1450	0.1531± 0.0037	0.1476	0.1335± 0.0028	0.1296
	0.0063	SAR	i		0.0037	SKCM	0.0028	
Miss-	TDi		TDi		 	SKCIVI	TDir	
ing	pute-DNA		pute-DN		TDimpute-DNA	Ameth_self	pute-DN	
rate	cv(5fold)	test	cv(5fold)	test	cv(5fold)	test	cv(5fold)	test
20%	0.1900 <u>±</u> 0.0066	0.1849	0.1790± 0.0063	0.1768	0.1638± 0.0037	0.1637	0.1582± 0.0035	0.1590
40%	0.1953± 0.0066	0.1906	0.1832± 0.0063	0.1810	0.1702± 0.0039	0.1704	0.1627± 0.0036	0.1634
60%	0.2036± 0.0069	0.1984	0.1895± 0.0062	0.1872	 0.1780± 0.0043	0.1783	0.1682± 0.0037	0.1692
80%	0.2236±	0.2168	0.2029±	0.1992	 0.1935±	0.1923	0.1790±	0.1790
	0.0081	STAI	0.0071		 0.0050	THCA	0.0040	
Miss-	TDi		TDi		 	INCA	TDir	
ing	pute-DNA		pute-DN		TDimpute-DNA	Ameth_self	pute-DN	
rate	cv(5fold)	test	cv(5fold)	,	 cv(5fold)	test	cv(5fold)	test
20%	0.1379± 0.0031	0.1376	0.1278± 0.0028	0.1290	 0.1278± 0.0036	0.1277	0.1267± 0.0031	0.1262
40%	0.1441± 0.0030	0.1442	0.1316± 0.0030	0.1328	0.1309± 0.0036	0.1326	0.1293± 0.0032	0.1286
60%	0.1527± 0.0034	0.1533	0.1370± 0.0028	0.1380	 0.1362± 0.0036	0.1416	0.1324± 0.0032	0.1318
80%	0.1699± 0.0043	0.1694	0.1475± 0.0031	0.1478	0.1482± 0.0037	0.1532	0.1374± 0.0035	0.1364

Table	Table S2 : The RMSE of five methods on 16 cancer datasets. Optimal results are highlighted in bold.												
Missi				BLCA			BRCA						
ng rate	Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h		Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h		
20%	0.23 0	0.1 94	0.1 67	0.156	0.143		0.21 6	0.1 80	0.1 55	0.144	0.139		
40%	0.23 1	0.2 31	0.1 68	0.161	0.145		0.21 6	0.2 15	0.1 55	0.149	0.141		

T	T		T			1				
CO0/	0.23	0.2	0.1	0.400	0.1.10	0.21	0.2	0.1	0.454	0.4.5
60%	1	61	70	0.168	0.149	7	48	56	0.154	0.145
0.00/	0.23	0.2	0.1			0.21	0.2	0.1		
80%	1	87	79	0.179	0.156	7	87	61	0.158	0.153
Missi		4		CESC	-		•		COAD	-
ng	Me	KN	SV	TDim-	TDim-	Me	KN	SV	TDim-	TDim-
_	an	N	D	pute-DNAmeth_s	pute-DNAmet	an	N	D	pute-DNAmeth_s	pute-DNAmet
rate				elf	h	all	11	ע	elf	h
	0.22	0.2	0.1	0.158	0.149	0.19	0.1	0.1	0.136	0.129
20%	3	01	70	0.130	0.143	6	63	41	0.100	0.123
	0.22	0.2	0.1			0.19	0.1	0.1		
40%	4	36	71	0.164	0.151	6	96	42	0.140	0.131
	0.22	0.2	0.1			0.19	0.2	0.1		
60%	4	67	75	0.171	0.155	6	24	45	0.145	0.135
	0.22	0.3	0.1			0.19	0.2	0.1		
80%	5	09	89	0.183	0.163	7	51	53	0.151	0.142
Missi				HNSC					KIRC	
	3.6	773.7	GT.	TDim-	TDim-	3.6	173.1	GT.	TDim-	TDim-
ng	Me	KN	SV	pute-DNAmeth s	pute-DNAmet	Me	KN	SV	pute-DNAmeth s	pute-DNAmet
rate	an	N	D	elf	h	an	N	D	elf	h
	0.19	0.1	0.1	0.400		0.16	0.1	0.1	0.110	
20%	4	61	37	0.128	0.120	1	29	12	0.113	0.108
	0.19	0.1	0.1			0.16	0.1	0.1		
40%	4	99	37	0.132	0.122	1	43	13	0.116	0.110
	0.19	0.2	0.1			0.16	0.1	0.1		
60%	4	40	39	0.137	0.125	1	74	15	0.119	0.113
	0.19	0.2	0.1			0.16	0.2	0.1		
80%	5	75	44	0.146	0.131	2	14	24	0.121	0.118
Missi		4	4	KIRP				4	LGG	1
				TDim-	TDim-				TDim-	TDim-
ng	Me	KN	SV	pute-DNAmeth s	pute-DNAmet	Me	KN	SV	pute-DNAmeth s	pute-DNAmet
rate	an	N	D	elf	h	an	N	D	elf	h
	0.19	0.1	0.1			0.18	0.1	0.1		•
20%	5	69	41	0.134	0.128	3	61	29	0.126	0.123
	0.19	0.1	0.1			0.18	0.1	0.1		
40%	5	93	42	0.137	0.131	3	90	30	0.130	0.125
	0.19	0.2	0.1			0.18	0.2	0.1		
60%	6	19	46	0.142	0.134	3	45	32	0.134	0.129
	0.19	0.2	0.1			0.18	0.2	0.1		
80%	6	63	60	0.153	0.141	3	71	41	0.141	0.135
N 4' '		i	1	LIHC	<u>i</u>		<u> </u>	i	LUAD	<u> </u>
Missi			<u> </u>	TDim-	TDim-			<u> </u>	TDim-	TDim-
ng	Me	KN	SV	pute-DNAmeth s	pute-DNAmet	Me	KN	SV	pute-DNAmeth s	pute-DNAmet
rate	an	N	D	elf	h	an	N	D	elf	h
	0.22	0.1	0.1			0.16	0.1	0.1		
20%	6	86	58	0.145	0.142	8	45	22	0.115	0.106
	0.22	0.2	0.1			0.16	0.1	0.1		
40%	7	33	59	0.149	0.145	8	83	23	0.119	0.108
	0.22	0.2	0.1	0.110	3.2.0	0.16	0.2	0.1	0.110	0.200
60%	7	85	61	0.154	0.148	8	13	25	0.124	0.111
	0.22	0.3	0.1	0.101	3.2.0	0.16	0.2	0.1	V.12 /	V.222
80%	7	11	69	0.163	0.156	9	41	30	0.131	0.117
	<u>'</u>	i**	1	LUSC	3.100	<i>J</i>	1 71	1	PRAD	1 0.111
Missi		<u> </u>	1	·	TD.			<u> </u>	***************************************	TD.
ng	Me	KN	SV	TDim-	TDim-	Me	KN	SV	TDim-	TDim-
rate	an	N	D	pute-DNAmeth_s	pute-DNAmet	an	N	D	pute-DNAmeth_s	pute-DNAmet
<u> </u>	<u> </u>	<u> </u>		elf	h	<u>]</u>	<u> </u>	<u> </u>	elf	h

	0.20	0.1	0.1	0.132	0.122	0.18	0.1	0.1	0.112	0.109
20%	4	71	44	0.102	0.122	0	52	20	0.112	0.100
	0.20	0.2	0.1			0.18	0.1	0.1		
40%	4	01	45	0.137	0.124	0	85	20	0.117	0.111
	0.20	0.2	0.1			0.18	0.2	0.1		
60%	5	46	47	0.143	0.127	0	12	22	0.121	0.114
	0.20	0.2	0.1			0.18	0.2	0.1		
80%	5	79	56	0.154	0.133	0	58	30	0.130	0.121
Missi				SARC					SKCM	
ng	Me	KN	SV	TDim-	TDim-	Me	KN	SV	TDim-	TDim-
rate	an	N	D	pute-DNAmeth_s	pute-DNAmet	an	N	D	pute-DNAmeth_s	pute-DNAmet
Tate		11	ט	elf	h	an		ט	elf	h
	0.25	0.2	0.1	0.184	0.175	0.23	0.2	0.1	0.156	0.156
20%	2	21	93	0.104	0.173	4	05	72	0.130	0.130
	0.25	0.2	0.1			0.23	0.2	0.1		
40%	2	59	95	0.189	0.178	4	42	73	0.161	0.158
	0.25	0.3	0.2			0.23	0.2	0.1		
60%	3	00	00	0.196	0.183	4	72	75	0.168	0.163
	0.25	0.3	0.2			0.23	0.3	0.1		
80%	4	31	18	0.211	0.193	5	04	82	0.180	0.172
Missi			_	STAD					THCA	
ng	Me	KN	SV	TDim-	TDim-	Me	KN	SV	TDim-	TDim-
rate	an	N	D	pute-DNAmeth_s	pute-DNAmet	an	N	D	pute-DNAmeth_s	pute-DNAmet
Tate		11	ט	elf	h		11		elf	h
	0.20	0.1	0.1	0.134	0.125	0.15	0.1	0.1	0.127	0.124
20%	6	71	43	0.101	0.120	 6	43	27	0.121	0.12
400	0.20	0.2	0.1			0.15	0.1	0.1		
40%	6	20	44	0.139	0.127	6	53	28	0.129	0.126
0.004	0.20	0.2	0.1			0.15	0.1	0.1		
60%	6	62	46	0.144	0.131	6	68	29	0.131	0.129
	0.20	0.2	0.1			0.15	0.1	0.1		
80%	6	88	55	0.154	0.138	6	95	33	0.135	0.134

Table S.	3 : The	R^2 of 1	five me	thods on 16 cancer of	datasets. Optimal	res	sults ar	re high	lighte	d in bold.	
Miss-				BLCA						BRCA	
ing rate	Me an	KN N	SV D	TDim- pute-DNAmeth_ self	TDim- pute-DNAmet h		Me an	KN N	SV D	TDim- pute-DNAmeth_ self	TDim- pute-DNAmet h
20%	0.8 66	0.9 05	0.92 8	0.940	0.947		0.8 75	0.9 11	0.9 32	0.944	0.948
40%	0.7 40	0.7 50	0.85 6	0.875	0.892		0.7 61	0.7 75	0.8 68	0.887	0.897
60%	0.6 18	0.5 69	0.77 9	0.797	0.832		0.6 51	0.6 08	0.8 03	0.820	0.840
80%	0.4 99	0.3 71	0.68 4	0.693	0.755		0.5 45	0.3 75	0.7 23	0.734	0.767
Miss-		i	i	CESC	.i			i	i	COAD	<u>i</u>
ing rate	Me an	KN N	SV D	TDim- pute-DNAmeth_ self	TDim- pute-DNAmet h		Me an	KN N	SV D	TDim- pute-DNAmeth_ self	TDim- pute-DNAmet h
20%	0.8 77	0.8 98	0.92 4	0.935	0.943		0.9 00	0.9 31	0.9 46	0.953	0.956
40%	0.7 65	0.7 53	0.85 2	0.867	0.889		0.8 08	0.8 19	0.8 95	0.904	0.913

	0.6	0.5	0.77			 0.7	0.6	0.8			
60%	58	79	3	0.785	0.828	21	74	41	0.847	0.865	
	0.5	0.3	0.66			 0.6	0.4	0.7			
80%	55	36	8	0.675	0.752	34	88	70	0.771	0.803	
Miss-		•		HNSC			•	-	KIRC	•	
ing	Me	KN	SV	TDim-	TDim-	Me	KN	SV	TDim-	TDim-	
rate	an	N	D D	pute-DNAmeth_	pute-DNAmet	an	N	D D	pute-DNAmeth_	pute-DNAmet	
Tate				self	h	 			self	h	
200/	0.8	0.9	0.94	0.952	0.957	0.9	0.9	0.9	0.972	0.974	
20%	93 0.7	25 0.7	4 0.88			 42 0.8	62 0.9	71 0.9			
40%	92	90	9	0.901	0.914	0.8 88	0.9	42	0.944	0.949	
4070	0.6	0.6	0.83	0.301	0.514	 0.8	0.8	0.9	0.344	0.343	
60%	96	02	2	0.841	0.866	36	18	11	0.912	0.921	
	0.6	0.3	0.76			 0.7	0.6	0.8			
80%	02	81	3	0.763	0.805	85	46	70	0.872	0.887	
Miss-		•	•	KIRP			•	•	LGG		
ing	Me	KN	SV	TDim-	TDim-	 Me	KN	SV	TDim-	TDim-	
rate	an	N	D D	pute-DNAmeth_	pute-DNAmet	an	N	D D	pute-DNAmeth_	pute-DNAmet	
Tate				self	h	 			self	h	
200/	0.9	0.9	0.96	0.966	0.969	0.9	0.9	0.9	0.970	0.971	
20%	28 0.8	46 0.8	2			 39 0.8	52 0.8	68 0.9			
40%	0.8 65	0.8 71	0.92 5	0.930	0.939	0.8 84	0.8 19	37	0.939	0.942	
7070	0.8	0.7	0.88	0.930	0.333	 0.8	0.6	0.9	0.939	0.342	
60%	0.0	71	4	0.888	0.904	34	54	0.3	0.905	0.910	
	0.7	0.5	0.82			 0.7	0.5	0.8			
80%	51	83	3	0.830	0.861	87	11	53	0.862	0.868	
Miss-		•	•	LIHC		LUAD					
ing	Me	KN	SV	TDim-	TDim-	Me	KN	SV	TDim-	TDim-	
rate	an	N	D	pute-DNAmeth_	pute-DNAmet	an	N	D	pute-DNAmeth_	pute-DNAmet	
Tate				self	h	 			self	h	
20%	0.8 84	0.9 20	0.94	0.948	0.951	0.8 94	0.9 20	0.9 40	0.950	0.954	
ZU70	0.7	0.7	0 0.88			 0.7	0.7	0.8			
40%	79	80	3	0.894	0.902	99	76	84	0.897	0.911	
1070	0.6	0.5	0.82	0.004	0.002	 0.7	0.5	0.8	0.001	0.011	
60%	81	86	2	0.831	0.848	07	97	24	0.835	0.862	
	0.5	0.4	0.74			 0.6	0.3	0.7			
80%	87	21	5	0.749	0.779	20	94	48	0.752	0.801	
Miss-				LUSC					PRAD		
ing	Me	KN	SV	TDim-	TDim-	Me	KN	SV	TDim-	TDim-	
rate	an	N	D	pute-DNAmeth_	pute-DNAmet	an	N	D	pute-DNAmeth_	pute-DNAmet	
Tate				self	h	 			self	h	
20%	0.8	0.9	0.93	0.949	0.955	0.9	0.9	0.9	0.970	0.971	
ZU%	85 0.7	17 0.7	9 0.87			 27 0.8	46 0.8	65 0.9			
40%	0.7 79	92	9	0.893	0.912	0.8 59	0.8 57	31	0.937	0.942	
. 570	0.6	0.5	0.81	0.000	0.012	 0.7	0.7	0.8	0.001	0.042	
60%	75	88	4	0.826	0.862	95	43	95	0.900	0.908	
	0.5	0.3	0.73			0.7	0.5	0.8			
80%	76	65	2	0.736	0.800	34	63	42	0.850	0.864	
Miss-				SARC					SKCM		
ing	Me	KN	SV	TDim-	TDim-	Me	KN	SV	TDim-	TDim-	
rate	an	N N	D D	pute-DNAmeth_	pute-DNAmet	an	N N	D D	pute-DNAmeth_	pute-DNAmet	
Tate	uII	1,	ע	self	h	 uii	1,	<u> </u>	self	h	

20%	0.8 81	0.9 08	0.92 7	0.936	0.942	 0.8 87	0.9 13	0.9 36	0.948	0.951
	0.7	0.7	0.85			 0.7	0.7	0.8		
40%	69	71	5	0.868	0.883	84	85	75	0.892	0.901
	0.6	0.5	0.77			0.6	0.6	0.8		
60%	62	89	4	0.789	0.818	86	36	11	0.826	0.846
	0.5	0.3	0.66			0.5	0.4	0.7		
80%	57	90	2	0.679	0.734	92	49	31	0.737	0.773
Miss-				STAD					THCA	
ing rate	Me an	KN N	SV D	TDim- pute-DNAmeth_ self	TDim- pute-DNAmet h	Me an	KN N	SV D	TDim- pute-DNAmeth_ self	TDim- pute-DNAmet h
20%	0.8 72	0.9 11	0.93 5	0.945	0.951	0.9 58	0.9 65	0.9 72	0.972	0.973
•	~ ~				•	 		!		
	0.7	0.7	0.87			0.9	0.9	0.9		
40%	0. <i>7</i> 57	0.7 39	0.87 2	0.887	0.903	0.9 18	0.9 22	0.9 44	0.943	0.946
40%				0.887	0.903	 			0.943	0.946
40%	57	39	2	0.887 0.817	0.903 0.849	 18	22	44	0.943 0.912	0.946
	57 0.6	39 0.5	2 0.80			 18	22 0.8	44 0.9		

	D4. 111	c c m	40/1 01		cancer datasets. O	F		ourto u		mgmea m cora.	
Missi				BLCA						BRCA	
ng rate	Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h		Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h
20%	0.59 1	0.5 96	0.5 99	0.59	0.593		0.61 9	0.6 21	0.6 27	0.625	0.625
40%	0.53 4	0.5 67	0.5 82	0.574	0.614		0.52 6	0.5 19	0.6 17	0.616	0.606
60%	0.51 4	0.5 31	0.5 42	0.591	0.614		0.51 6	0.5 42	0.5 84	0.603	0.615
80%	0.49 4	0.5 08	0.5 75	0.589	0.575		0.54 7	0.5 16	0.5 78	0.595	0.603
Missi	CESC						COAD				
ng rate	Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h		Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h
20%	0.65 7	0.6 59	0.6 77	0.688	0.693		0.52 7	0.5 32	0.5 4	0.516	0.555
40%	0.63 7	0.6 12	0.6 28	0.668	0.642		0.56 7	0.6 03	0.5 03	0.528	0.564
60%	0.51 3	0.5 42	0.6 01	0.616	0.598		0.57 4	0.5 65	0.5 44	0.58	0.6
80%	0.47 3	0.5 16	0.6 08	0.61	0.596		0.50 8	0.4 69	0.5 1	0.515	0.55
Missi	HNSC									KIRC	
ng rate	Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h		Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h
20%	0.59 3	0.6 07	0.6 09	0.613	0.611		0.68 7	0.6 94	0.6 89	0.696	0.693
40%	0.60	0.6	0.6	0.616	0.62		0.68	0.6	0.6	0.682	0.687

	2	2	28			8	79	86		
60%	0.51	0.5 46	0.5 78	0.593	0.601	0.69	0.6 98	0.6 96	0.697	0.701
80%	0.48 9	0.5 01	0.5 41	0.55	0.563	0.66	0.6 64	0.7 03	0.717	0.728
Missi		i	i	KIRP			i	i	LGG	<u>i</u>
ng rate	Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h	Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h
20%	0.82 7	0.8 23	0.8 25	0.824	0.821	0.82	0.8 26	0.8 29	0.827	0.829
40%	0.79 1	0.8 11	0.8 3	0.814	0.824	0.80	0.8 06	0.8 28	0.823	0.832
60%	0.68	0.6 89	0.8 15	0.808	0.788	0.79 8	0.7 78	0.8 27	0.82	0.833
80%	0.65	0.7 33	0.8 15	0.801	0.803	0.76 1	0.7 35	0.8 2	0.819	0.837
Missi				LIHC					LUAD	<u> </u>
ng rate	Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h	Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h
20%	0.57 9	0.5 98	0.6 03	0.586	0.581	0.61 4	0.6 03	0.5 91	0.599	0.606
40%	0.51	0.5 54	0.5 58	0.563	0.565	0.62 4	0.6 14	0.6 05	0.608	0.608
60%	0.58 6	0.5 8	0.5 96	0.56	0.546	0.51 1	0.5 38	0.5 58	0.591	0.618
80%	0.53 6	0.5 33	0.5 78	0.541	0.544	0.54 3	0.5 63	0.5 28	0.53	0.595
Missi				LUSC					PRAD	
ng rate	Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h	Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h
20%	0.55 8	0.5 51	0.5 46	0.544	0.562	0.48	0.5 54	0.5 58	0.596	0.602
40%	0.52 8	0.5 54	0.5 43	0.58	0.541	0.5	0.5	0.5	0.5	0.5
60%	0.54 6	0.4 98	0.5 67	0.563	0.56	0.5	0.5	0.5	0.5	0.5
80%	0.49 9	0.4 95	0.5 79	0.558	0.581	0.5	0.5	0.5	0.5	0.5
Missi			·····	SARC				•	SKCM	•
ng rate	Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h	Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h
20%	0.68	0.6 91	0.6 68	0.682	0.673	0.6 61	0.6 53	0.6 51	0.645	0.645
40%	0.69 3	0.6 94	0.6 68	0.677	0.672	0.64	0.6 36	0.6 19	0.638	0.655
60%	0.53 8	0.5 44	0.6 38	0.636	0.645	0.51	0.5 12	0.6 08	0.619	0.638
80%	0.56 9	0.5 87	0.6 43	0.633	0.645	0.50	0.5	0.5 57	0.608	0.625
Missi		i	4	STAD			i	i	THCA	4

Imputing DNA methylation by transferred learning

ng rate	Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h	Me an	KN N	SV D	TDim- pute-DNAmeth_s elf	TDim- pute-DNAmet h
20%	0.5	0.5	0.5	0.5	0.5	0.79 7	0.7 96	0.7 95	0.791	0.801
40%	0.5	0.5	0.5	0.5	0.5	0.79 9	0.8 07	0.8 06	0.801	0.808
60%	0.5	0.5	0.5	0.5	0.5	0.65 8	0.7 54	0.7 64	0.793	0.764
80%	0.5	0.5	0.5	0.5	0.5	0.62 7	0.7 25	0.7 05	0.782	0.784