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28 Abstract

Type 2 Diabetes (T2D) diagnosis is based solely on glycemia, even though it is an endpoint of
numerous dysmetabolic pathways. T2D complexity is challenging in a real-world scenario, thus
dissecting T2D heterogeneity is a priority. Cluster analysis, which identifies natural clusters
within multidimensional data based on similarity measures, poses as a promising tool to unravel
Diabetes complexity.

Herein, we aimed at scrutinizing and integrate the results obtained in most of the works up to date. We conclude that to correctly stratify subjects and to differentiate and individualize a preventive or therapeutic approach to Diabetes management, cluster analysis should be informed with more parameters than the traditional ones, such as etiological factors, pathophysiological mechanisms, other dysmetabolic co-morbidities, and biochemical factors i.e. the *millieu*. Ultimately the abovementioned factors may impact on Diabetes and its complications.

Lastly, we propose another theoretical model, which we named the Integrative Model. We
differentiate three types of components: etiological factors, mechanisms, and millieu. Each
component encompasses several factors to be projected in separate 2D planes allowing an holistic
interpretation of the individual pathology.

Fully profiling the individuals, considering genomic and environmental factors, and exposuretime, will allow the drive to precision medicine and prevention of complications.

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47 Keywords: diabetes; machine learning; cluster analysis; big data

49	Abbreviations
50	BMI – body mass index
51	CAD – coronary artery disease
52	CKD – chronic kidney disease
53	CV – cardiovascular
54	DKD – diabetic kidney disease
55	eGFR – estimated glomerular filtration rate
56	GRS – genetic risk score
57	HOMA-B – Homeostatic Model Assessment for beta-cell function
58	HOMA-IR – Homeostatic Model Assessment for Insulin Resistance
59	MARD – mild-age related Diabetes
60	ML – machine learning
61	MOD – mild-obesity related Diabetes
62	MR – Mendelian Randomisation
63	MRI – magnetic resonance imaging
64	NAFLD – non-alcoholic fatty liver disease
65	OAD – oral antidiabetic drugs
66	OGTT – oral glucose tolerance test
67	PAM – partition around medoids
68	PD – Prediabetes
69	SAID – severe autoimmune Diabetes
70	SIDD – severe insulin-deficient Diabetes
71	SIRD – severe insulin-resistant Diabetes
72	SNPs – single nucleotide polymorphisms
73	SOM – self organizing maps
74	T1D – Type 1 Diabetes mellitus
75	T2D – Type 2 Diabetes mellitus
76	UACR – urine albumin creatinine ratio

77 1. Introduction

78 In Diabetes glucose metabolism is affected due to individual or simultaneous changes in insulin 79 secretion, action or metabolism. Diabetes is diagnosed based on glycemia and cut-off values were defined based on the presence of microvascular complications, namely retinopathy.¹ However, 80 dysglycemia, or the glucose altered metabolism, is not an all-or-nothing phenomenon on the 81 82 contrary, it occurs continuously. Prediabetes (PD) is a less severe hyperglycemic state that depicts 83 a higher risk of progression to Diabetes. Importantly, individuals with PD can develop Diabetes 84 complications, whereas others with Diabetes may never develop them, showing the limitations of the current clinical classification.² Therefore, glycemic levels are not sufficient to inform about 85 86 the onset and severity of the condition.

87 Notwithstanding all investment in Diabetes, specifically in Type 2 Diabetes mellitus (T2D), it is 88 still one of the main non-communicable diseases, and its mortality increased 70% since 2000.³ T2D is extremely heterogenous,^{4,5} both in its initial presentation and complications' development, 89 90 which is crucial to explain the sustained morbidity and increased mortality attributable to this condition.^{3,6} The empirical individualisation of therapy in Diabetes dates back to 19th century,^{7,8} 91 92 and is still practised. The latest therapeutic guidelines for T2D include several recent drugs that are giving better results regarding cardiometabolic complications⁹ and start to have an increased 93 94 focus on the patient's co-morbidities.¹⁰ The concept of precision medicine has been proposed, 95 aiming at defining the most effective approach for a similar group of patients regarding genetic, environmental, lifestyle, clinical factors, amongst others.⁶ However, further advances in the 96 97 ability to define precise therapies for Diabetes also depend on the acquired knowledge regarding 98 the heterogenity of the condition.

As early as 1965, two major groups were acknowledged in Diabetes pathophysiology: insulin resistant and insulin deficient individuals.¹¹ The two pathophysiological mechanisms associated with these groups were assumed to be related with two main organs: insulin secretion impairment in the pancreas; and insulin resistance at the skeletal muscle. Since then, much more complexity was added to Diabetes pathophysiology, especially to T2D.¹² More recently, it has been shown that other organs and factors, such as the lung and microbiome, can impact on T2D onset and progression.¹³⁻¹⁵ Additionally, it is currently accepted that T2D etiology encompasses thousands
of low impactful genes, as well as environmental and lifestyle factors, that interact with each
other.¹⁶

108 Glucose metabolism is part of an intricate metabolic network where carbohydrates, lipids and 109 other metabolic pathways should be considered as a whole and, when affected, result in 110 dysmetabolism and/or hemodynamic alterations. Thus, depending on the affected mechanisms, 111 Diabetes can appear in distinct dysmetabolic contexts. Interestingly, there are lipodystrophic 112 phenotypes in which the inability of white adipose tissue to expand, despite diverse BMI values, causes ectopic fat deposition.¹⁷ These subjects are exposed to atherogenic dyslipidemia¹⁸ and, in 113 114 the liver, development of fatty liver may progress to steatohepatitis¹⁹ that can be further impacted 115 by different adipose tissue amounts and function. Despite showing similar patterns regarding 116 hyperglycemia and hyperlipidemia, subjects with lipodystrophy, might require a distinct treatment.²⁰ Another example relates to Diabetes and hypertension bidirectional association. Both 117 118 conditions have several common pathophysiological mechanisms, namely hyperinsulinemia, 119 increased sympathetic nervous activity, activation of renin-angiotensin-aldosterone system, endothelial dysfunction, etc.²¹ The onset of hypertension in subjects with Type 1 Diabetes (T1D) 120 121 has been related with the onset of kidney dysfunction; however, in subjects with T2D, it can appear before²² and they can show a prehypertensive profile some years earlier.²³ The causal 122 123 association of T2D in hypertension was depicted in a Mendelian Randomisation (MR) study, but does not explain the onset of T2D in hypertensive subjects.²⁴ However, a higher incidence of T2D 124 in hypertensive subjects as compared with normotensive subjects is evident.²⁴ The above-125 126 described complexity, although easy to understand in concept, is very hard to demonstrate and 127 tackle in clinical practice. Dissecting and understanding T2D heterogeneity is a priority to reverse the current scenario.25 128

129 To tackle the overly complex clinical challenges, involving multiple etiological factors, organs 130 and mechanisms, classical statistical analyses are frankly insufficient. Recent progress in memory 131 and computation power allowed for the development and implementation of more complex algorithms, including a collection of tools that can learn from data, named machine learning (ML).
Specifically cluster analysis, using unsupervised learning algorithms (algorithms that deal with
observations that do not have a label to learn from²⁶) are promising tools to unravel Diabetes
complexity.

We will critically review distinct cluster analysis methodologies currently used to study Diabetes and integrate results from different studies. Since all analyses aimed at understanding Diabetes/T2D pathophysiology, we anticipate their conclusions to fit as pieces on a puzzle. Finally, we suggest a model that can be applied to Diabetes precision medicine and from a wider perspective to dysmetabolism overall.

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142 2. Advancement of Diabetes Management – travelling on the road to precision medicine

143 The word Diabetes ("to go through" or siphon) is attributed to Apollonius of Memphis in Greece 144 around 250BC. However, its clinical description and some complications date back to 3500 years ago in Egypt.²⁷ Interestingly, two types of Diabetes - congenital and late onset - and their 145 146 relationship to heredity, obesity, sedentariness and diet, were already recognized in medical treatments in ancient India.^{8,28} At the time Diabetes resulted in death and preventing it was the 147 148 main goal. Additionally, complications of Diabetes, as peripheral neuropathy, gangrene and erectile dysfunction were described by an Arab doctor, Avicenna (AD 960-1037).²⁷ Centuries 149 150 later Matthew Dobson (1732-1784) and Michel Chevreul (1786-1889), through the application of 151 chemistry to diagnosis, identified glucose as the sugar that was increased both in urine and serum of these patients.⁸ Arguing that glucose appeared in the urine because the body was unable to 152 153 assimilate it, Dobson considered Diabetes a systemic disease rather than a kidney disease, as it was considered until then.28 These findings led to the research on the metabolism of 154 155 carbohydrates. However, insulin was not yet available and treatments were based on individualisation of diets, rest or other lifestyle changes,⁷ unable to prevent death from acute 156 157 complications. Neurological complications were also quite frequent, the association of 158 neuropathy, vascular disease, plantar ulcers and gangrene with Diabetes was also described, rising the hypothesis that microvascular disease was the cause of some complications.²⁸ 159

160 In 1921-22 Banting and Best isolated insulin, one of the great discoveries in medicine, which has 161 allowed most people with insulin-dependent Diabetes to be treated to this day. On the other hand 162 it led to the distinction of T1D, in which people needed insulin, from T2D, in which insulin was present but uneffective.²⁷ Since the problem in question was hyperglycemia, other therapeutic 163 strategies would be developed based on glycemic control.²⁷ In the 1950's the first sulfonylurea 164 appeared - the first oral antidiabetic drug (OAD) for people with T2D.²⁹ Metformin, the most used 165 OAD, appeared a few years later with its mechanism of action only recently fully understood.³⁰ 166 167 Since then, other groups have been made available as the involvement of other organs and mechanisms is known.^{10,12,29} In a paradigm of therapy which in the meantime has become 168 169 evidence-based clinical guidelines began to be published, with the main therapeutic focus on glycemic control.³¹ It was also recognized that the reduction of complications implied 170 171 simultaneous treatment of other diseases that represent risk factors for the same complications, 172 such as dyslipidemia and hypertension.³¹

The etiologic classification of Diabetes recognizes several types besides Type 1, Type 2 and gestational Diabetes.¹ The recognition that there is still a high degree of heterogeneity leads to an effort to adapt the numerous drugs with distinct mechanisms to the patients who benefit most from them.³² Weight control, hypertension and dyslipidemia, among others, have gained increasing relevance along with glycemic control.¹⁰ Nowadays, these diseases are recognized as co-morbidities but treated as independent conditions.

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180 3. Cluster Analysis

181 Cluster analysis is a ML methodology that uses a group of algorithms that can deal with non-182 labelled data, named unsupervised learning (Figure 1). Cluster analysis aims to stratify population 183 observations' in natural groups/clusters without needing *a priori* categorization. Within each 184 cluster observations' similarity are maximized whilst minimized between clusters.³³

185 Distinct clustering algorithms have advantages and drawbacks related to computation time, the 186 need for an *a priori* knowledge regarding the number of groups, and cluster shape in a 187 multidimensionality space that they can find (Table 1).²⁶ In (dys)glycemia, specifically in the resolution of T2D heterogeneity, one should consider several parameters with distinct and specific
characteristics (e.g. genes, environmental factors, biochemical analysis, omics, etc.). Therefore,
it is natural that the best result is obtained using an ensemble of algorithms.

191 Cluster analysis workflow implies taking several decisions (e.g. choosing the algorithm, variables 192 to inform the cluster, similarity and distance measures, etc.). When algorithms are not able to find 193 the best number of clusters (Table 2), there is the need to determine *a priori* a number of clusters.³⁴ 194 Still, different measures can give a distinct optimal number of clusters and therefore should be 195 carefully selected and interpreted. Of note, the found groups should be clinically relevant. 196 Furthermore, aside from finding natural groups in data, cluster analysis is a powerful tool in data 197 exploration and visualization. In the context of (dys)glycemia heterogeneity, by profiling the 198 found groups, we can explore what characterizes them, posing a promising tool to explore and 199 tackle (dys)glycemia complexity.

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201 4. Cluster Analysis Algorithm impact on Founded Clusters

To perform a cluster analysis, impactful decisions must be made: inclusion and exclusion criteria, choice of variables, and the algorithm to perform the analysis, amongs others. Additionally, indexes that define the best number of clusters and distance metrics have to be selected.²⁶ Cluster analysis used to date to tackle T2D and dysmetabolism have a dissimilar methodology that must be considered when interpreting and integrating the results (Table 2).^{35–38}

207 Hierarchical clustering and k-means are two of the most well-known clustering algorithms. Agglomerative hierarchical clustering²⁶ is a simple algorithm that hierarchically joins nested 208 209 clusters in a bottom-up way, with its agglomerative process visualized in a dendrogram. This 210 process does not need the pre-specification of the optimal number of clusters, though it requires 211 an *a posteriori* cut-off to define them. Furthermore, data can be analyzed at different cut-off 212 values, allowing us to understand how observations aggregate. However it can only find clusters 213 with specific shapes, it gives distinct solutions depending on the chosen aggregation methodology to join the observations and has a high computation cost.²⁶ K-means is a simple and efficient 214 algorithm. Besides not dealing well with categorical variables, the final solution is highly 215

impacted by its random initialization, requires an *a priori* specification of the number of clusters and, importantly, it is prone to find spherical clusters, even if this is not their natural shape.²⁶ The latter can limit its use. Partition around medoids (PAM) is a *k*-medoids algorithm, that is less sensitive to noise than *K*-means, but with a higher computational cost.²⁶

K-means, PAM and hierarchical clustering have been used mainly when few parameters are used
 to tackle T2D.³⁹ To perform more complex analyses, self-organising maps (SOMs) and
 topological based analysis have proven to be more efficient and able to find clusters that have
 non-spherical shapes.^{26,40}

Hierarchical SOMs, followed by hierarchical clustering,⁴¹ have been used to solve multiple 224 intricate problems, including clustering analysis of T1D complications.⁴⁰ SOM is a neural 225 226 network-based algorithm, which maps observations to neurons in a grid that at the end will represent the cluster (cluster centroid).⁴² In summary, the first algorithm allows data 227 228 dimensionality reduction, whereas the second enables the stratification and understanding of how 229 the units agglomerate together. Aside from dealing with large and complex data, SOMs can find 230 different cluster formats. Nonetheless, it has drawbacks as requiring too many parameters to be set and optimised, its computational cost and the number of clusters must be set a priori.⁴² 231 232 Network analysis is a graph-based method that assesses subjects (nodes) in relation to each other (edges).³⁶ 233

The abovementioned algorithms are classified as hard clustering algorithms, i.e. they group the population to assign one subject only to one cluster. Contrarily, soft clustering uses algorithms that define the probability of one observation belonging to distinct clusters;^{43,44} thus, one subject can belong to multiple clusters at a given time. Despite computational cost and convergence drawbacks, soft clustering algorithms are extremely useful when an item can belong to more than one cluster, as is the case of clustering T2D related genes/SNP's and mechanisms.³⁸

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241 5. Population and Parameter Set to resolve Type 2 Diabetes

Clusters analyses to resolve T2D heterogeneity are also diverse regarding the analyzed
population, set of parameters used to inform the cluster,^{40,43-45} thus impacting on the groups

found. Methodological heterogeneity reveals the authors' distinct perspectives on Diabetes
definition, where it stands within the wider dysmetabolism concept, and the number and type of
parameters that allows a precision medicine approach to T2D.

Although T2D is classically considered an affection of glucose metabolism, glucose metabolism 247 occurs integrated with other substrates'.⁴⁵ Glucose metabolism-related parameters though 248 informing about groups with different conditions, do not give a broader perspective on 249 250 metabolism nor account for the overall metabolic heterogeneity. T2D impact relies mainly on its complications' development that, in turn, relate to other factors.⁴⁶ Herein, we distinguish 251 252 etiological factors (e.g. time, genes, environmental factors, lifestyle factors), pathophysiological 253 mechanisms (e.g. overall or organ-specific insulin resistance, insulin secretion, overall or organ-254 specific insulin clearance), other dysmetabolic co-morbidities (e.g. hypertension, dyslipidemia), 255 biochemical and other internal environment factors present in the organism or that the organism 256 is exposed to, that is its *millieu* (e.g. glycemia, insulinemia, free fatty acids, blood pressure, body 257 weight).

258 Ahlqvist et al. performed a cluster analysis on a population of individuals recently diagnosed with Diabetes.³⁵ The analysis considered six clinical parameters: the presence of GAD antibodies 259 260 (GADA), age at diagnosis, HbA1c, BMI, HOMA-IR and HOMA-B. The analysis does not rely 261 only on glycemia nor on insulin levels. Nevertheless, the population solely includes individuals 262 that were diagnosed based on current criteria. The authors found five optimal clusters using the 263 silhouette index and hierarchical clustering. One of these clusters, named severe autoimmune 264 Diabetes (SAID), included GADA+ subjects. Afterwards, GADA+ subjects were excluded and 265 the k-means algorithm was used to define the other 4 clusters: severe insulin-deficient Diabetes 266 (SIDD); severe insulin-resistant Diabetes (SIRD); mild-obesity related Diabetes (MOD); and 267 mild-age related Diabetes (MARD). These clusters were replicated in other northern European cohorts.³⁵ In brief, SAID subjects showed an early-onset condition, low BMI and poor metabolic 268 269 control. Subjects in SIDD cluster were similar to SAID but GADA-; these subjects showed a 270 higher risk of having diabetic retinopathy. A variant in human leukocyte antigen (HLA) locus 271 (rs2854275) was found to be associated with SAID but not with SIDD.

272 Interestingly, Zaharia et al. showed that, in a German population, individuals that were GADA-273 at baseline could be GADA+ after five years, determining that for better classification of autoimmune Diabetes other antibodies should be used.⁴⁷ SIRD cluster included individuals with 274 275 marked insulin resistance, high BMI and a high prevalence of non-alcoholic fatty liver disease 276 (NAFLD). Of note, this cluster also revealed to have the highest β -cell function. Additionally, 277 individuals in the SIRD cluster were at the highest risk of developing chronic kidney disease 278 (CKD) and diabetic kidney disease (DKD, defined by persistent macroalbuminuria), despite 279 proper glycemic control. Finally, subjects in MOD showed higher values of BMI but not insulin 280 resistance, whereas MARD subjects were older, with only modest metabolic affection and were 281 not associated with the evaluated Diabetes complications. These last two clusters included most 282 of the population and still have a considerable proportion of subjects with Diabetes complications. 283 Furthermore, not all Diabetes' complications were evaluated. In fact, it has been suggested that 284 borderline Diabetes is associated with an increased risk of dementia and Alzheimer disease, which 285 is potentiated when hypertension is present. Regarding gene loci, rs7903146 (a TCF7L2 SNP 286 associated with T2D) was associated with SIDD, MOD and MARD; whereas rs10401969 (a TM6sf2 gene variant associated with NAFLD) was associated with SIRD but not with MOD.³⁵ 287 The above-mentioned four subgroups of T2D have been overall replicated, using the same 288 289 methodology as Ahlqvist et al., in distinct geographical locations and ethnicities. This further 290 confirms the already known association of Diabetes with younger subjects, with lower BMI and more insulin deficiency in Asian and Indian populations.^{48,49} Moreover, 23% of subjects changed 291 cluster in the five year follow-up.⁴⁷ Particularly, people in the insulin-deficient cluster (SIDD) 292 293 were changed to clusters with better insulin secretion (MOD and MARD).

Li *et al.* performed topology-based cluster analysis of 2552 T2D subjects from several ethnicities, informed by 73 mixed features from electronic medical records derived clinical data.³⁶ These features included biochemical and clinical parameters besides glycemia, thus approaching T2D in a wider (dys)metabolic context. This was a landmark study and one of the first studies to show the ability to deal with a high number of variables to stratify subjects with T2D. However, the stratification results depend on the parameters selected to inform the cluster rather than the chosen 300 population. It is not clear if the authors have found three Diabetes subtypes or three subtypes of 301 patients that have Diabetes, considering the highly mixed chosen parameters to inform the 302 analysis that also included several diseases codes and medications. The chosen methodology 303 renders it difficult to validate it in different populations.

304 To extend clusters' evaluation to subjects with normoglycemia and PD, we accounted for age as 305 a surrogate of time exposition, anthropometry, and biochemical parameters (glycemia, insulin, cpeptide and free fatty acids) in three-time points of an OGTT.³⁷ In this study, we used a 306 307 hierarchical SOM, followed by a hierarchical clustering algorithm. Subjects were then profiled 308 concerning the abovementioned parameters and several mechanism's surrogate indexes, including 309 overall and tissue-specific insulin resistance, insulin secretion, insulin clearance, NAFLD, and 310 glomerular filtration rate (GFR). The sample had a limited number of subjects with non-treated 311 T2D. Nonetheless, none of the subjects had Diabetes five years earlier. In this work, we found 312 two main clusters: one that includes subjects with a median metabolic phenotype compared to the 313 overall population; and the other with elevated insulin resistance and insulin secretion. However, 314 these 2 clusters were highly heterogeneous when they were evaluated for a higher number of 315 clusters. For example, despite the presence of a main insulin-resistant group, that comprised 316 subjects with normoglycemia and dysglycemia it included subgroups that could be differentiated 317 by their adipose tissue insulin resistance. Moreover, even though groups with lower estimated 318 GFR (eGFR) were insulin resistant, not all insulin resistant groups showed this association. 319 Additionally, we found that clusters including individuals with normo/dysglycemia and low 320 eGFR could be further profiled and showed insulin resistance and NAFLD. Consistently, other 321 groups have also shown that both high insulin resistance and NAFLD are related to kidney dysfunction in subjects with or without T2D.⁵⁰ In Ahlqvist et al. the group of individuals that had 322 323 the highest risk of developing CKD/DKD, even considering proper glycemic control, was the 324 most insulin resistant one.35

Furthermore, these subjects had the lowest GFR at baseline (when they had less than 12 months
from diagnosis) on the German Diabetes Study cohort.⁴⁷ The impact of insulin resistance and
NAFLD on GFR seems to be, at least partially, independent from glycemia. Importantly, both

328 conditions can be associated with hyperinsulinemia and insulin is a known trigger and a target of 329 kidney (dys)function that may have a role in the pathophysiology of T2D.⁵¹ Interestingly, the 330 heterogeneity of affected mechanisms was not exclusive of people with T2D, including also 331 subjects with PD and normoglycemia. Our work would benefit from being validated in other 332 cohorts. Nevertheless, we highlight that T2D diagnosis should consider other parameters besides 333 glycemia. In fact glucose levels impact is differently perceived by each individual. Therefore, it 334 should include subjects with different ranges of glycemic values together with other parameters.

335 An interesting complementary approach to dissect T2D heterogeneity is the use of genetic 336 markers. Reasoning that genetic variants remain constant despite disease progression and 337 treatment, unlike clinical variables, thus being more likely to reveal T2D causal mechanisms, a 338 cluster analysis including T2D gene-traits associations, including 94 genetic variants and 47 traits 339 was performed.³⁸ Aside from genetic data the analysis was informed with clinical parameters, 340 including surrogate indexes of insulin secretion and insulin resistance, as well as lipid parameters, 341 that allowed for the identification of other insulin resistance-related groups. Importantly, in this 342 work b-NMF, a soft clustering algorithm was used, allowing a SNP to be associated with more 343 than one mechanism and one cluster. The authors identified five clusters of genetic loci - traits 344 associations: two with variant-trait associations related to reduced β -cell function, distinct in pro-345 insulin levels; and three insulin resistance-related, namely obesity mediated, lipodystrophy-like 346 fat distribution and disrupted liver lipid metabolism. Of note, this is also a potentially complex 347 approach. As more than 100 loci were already found to be associated with T2D, each one with a 348 very slight impact on the increased risk of the disease and in dysmetabolism etiology, we should 349 consider, along with genetic factors, their interactions with environmental and lifestyle factors. 350 Interestingly, Udler et al. evaluated the Genetic Risk Score (GRS) association with relevant 351 outcomes in each cluster. Coronary artery disease (CAD) was mostly associated with the 352 lipodystrophy and Beta-cell clusters. Beta-cell cluster was also associated with ischemic stroke. 353 Increased blood pressure was only associated with lipodystrophy cluster, which also showed an 354 association with higher urine albumin-creatinine ratio (UACR). Liver/Lipid cluster was associated with decreased renal function and diminished UACR. GRS outcomes were validated

356 in T2D cohorts by profiling subjects' characteristics in top quantile GRS's subjects.³⁸

357 More recently, Wagner et al. focused on a german population considered at risk of developing Diabetes based on BMI, previous history and family history (TUEF/TULIP cohort).⁵² Besides 358 359 OGTT-based measures reflecting blood glucose, insulin resistance and insulin secretion, liver, 360 subcutaneous and visceral fat values measured by MRI, and HDL levels, polygenic risk score for 361 Diabetes were also included. The defined six clusters were then evaluated in a larger cohort 362 (Whitehall II). However to assign the latter individuals to the clusters, the authors used less 363 profiling variables, still based on OGTT measurements. The authors reported a relocation rate of 364 only 60% in the original cohort, which suggests that MRI fat measurements do not appear to be 365 superior to measurements such as BMI and waist circumference.³⁷ Importantly, progression to Diabetes, CKD, CV events and all cause mortality were assessed.⁵² Consistent with our findings,³⁷ 366 Wagner et al. demonstrated that pathophysiological affection is already present before Diabetes 367 diagnosis.⁵² Within the six defined clusters, three that were older and/or more obese showed 368 369 higher glycemia (clusters 3, 5 and 6); one related with insulin deficiency and raised genetic risk 370 (cluster 3); and two with insulin resistance (clusters 5 and 6). Cluster 6 showed a dissociation of 371 both risks of progression to Diabetes and CKD in Whitehall II cohort. However, considering that 372 GFR is not depicted in TULIP/TULIF and CKD progression models in Whitehall II were not 373 adjusted to GFR at the baseline these results should be carefully interpreted. Cluster 4 is consistent 374 with a metabolically health obese profile that includes younger subjects than the most 375 dysmetabolic groups and did not show a protected profile overtime, namely regarding CV events. 376 In fact, although clusters in TULIP/TULIF cohort differ in intima-media thickness, in the 377 Whitehall II cohort, the clusters did not differ in CV outcomes risk, after adjustment for BMI and 378 age, except for Cluster 2 that had a protected profile. Considering the relevance of CV events in Diabetes this highlights the importance of an enriched milieu to a better stratification.³⁷ 379

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381 6. What can we learn from cluster analysis?

382 Insulin secretion and resistance have been included in parameters informing cluster analyses. However there can be different mechanisms that lead to insulin deficiency and resistance.³⁷ It has 383 384 been suggested that insulin resistance can be considered a defensive mechanism against elevated insulin secretion due to a highly nutritional load in a sensitive β -cell.⁵³ In distinct cluster analysis, 385 386 most of the groups found to be insulin resistant were the ones with the highest insulin secretion.^{35,37} Nevertheless, the amount of circulating insulin depends not only on the cells' 387 secretion capacity but on overall insulin metabolism and on insulin clearance.⁵⁴ Changes in insulin 388 clearance have also been linked to hyperinsulinemia.^{37,54} Insulin resistance has been associated 389 390 with age and BMI. Interestingly, in work by Alqhvist et al., MARD and MOD groups differ from the SIRD in that they are less fat or younger, respectively, showing better metabolic control.³⁵ 391

392 Several questions remain to be clarified concerning the mechanisms leading to insulin resistance. 393 One concerns the mechanisms through which age and BMI impact on insulin resistance and 394 whether this implies a different therapeutic approach. Secondly, in the setting of insulin 395 resistance, it is known the association between liver and adipose tissue but whether insulin 396 resistance develops through distinct pathways, implying distinct therapeutic approaches, remains 397 elusive. Thirdly, when it comes to Diabetes complications, the majority of the results were 398 obtained using patients undergoing treatments, which may, in its turn, promote complication's onset.55 Finally, cluster analysis showed the association between GFR and albuminuria with 399 insulin-resistant states;^{40,45,52,56} however, the presence of an association does not necessarily imply 400 401 homogeneity between clusters, when it comes to kidney function, making this an etiological factor 402 of the uttermost importance in Diabetes stratification.

Udler using SNP and traits, in addition to HOMA-IR and HOMA-B, namely lipid profile, found
 three groups of insulin-resistant subjects that showed involvement of different mechanisms and
 organs.³⁸ We and others have shown that distinct insulin resistance patterns can be present in
 subjects with normoglycemia and PD.^{37,52}

407 Altogether these support the view that, in order to stratify subjects to differentiate a preventive or
408 therapeutic approach to Diabetes, one should inform the cluster analysis with more parameters
409 reflecting other mechanisms metabolites and factors (e.g., lipids, blood pressure, insulin).

Additionally, Diabetes pathophysiology occurs continuously and people without Diabetes can
already have Diabetes's complications, hinting to different susceptibilities to glycemic levels. This
may be due to concomitant exposure to other factors such as hypertension or dyslipidemia, or due
to the common underlying pathophysiologic mechanisms.

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415 7. New models for an approach to Diabetes in precision medicine

Cluster analysis is contributing to uncover the heterogeneity of Diabetes.^{35,37,38,47} However, its
superiority over simple predictive models (e.g., predicting complications such as renal
dysfunction) is being questioned.⁵⁶

McCarthy proposed the palette model to resolve T2D heterogeneity.57 The model defined 419 420 component planes, such as mechanisms, etiological factors and others, that can be affected, 421 comparing them to a palette hue. The characterization of subjects by their component planes 422 places them in a bidimensional plane where the path from normoglycemia to Diabetes can be 423 assessed for each individual. Importantly this model includes subjects with normoglycemia and 424 dysglycemia, which have different affected mechanisms. Ahlqvist et al. suggested a model based 425 on the assumption that there is a dominant pathway that gives at least to the majority of patients with Diabetes a well-defined "palette colour".58 Additionally, few clinical parameters render 426 427 larger groups.

428 In our view, a precision medicine model to approach Diabetes must consider glycemia and 429 glucose metabolism, as well as other substrates and factors, that impact on dysglycemia and/or 430 Diabetes complications onset and progression. Diabetes complications occur for different values 431 of glycemia, impacted by the metabolic context of the individual. In fact, dysmetabolic factors 432 interaction might potentiate the risk for specific conditions, as is the case of glycemia and blood pressure interaction in the development of Alzheimer's disease.⁵⁵ Finally, the model must be 433 434 holistic and applicable to different ethnicities. There are ethnicities that show a higher risk for the onset of T2D at younger ages and for lower BMI.⁴⁸ Interestingly, subjects with an Asian genetic 435 436 background seem to have diminished insulin secretory capacity, but one cannot exclude the 437 environmental and culture-related factors.

438 We propose to paint another picture, the Integrative model (Figure 2). We consider that the 439 approach can only be attained by being detailed in the metabolic characterization of the 440 individuals, and by placing it in a wider context of dysmetabolism. Thus, we consider the path 441 from normometabolism to dysmetabolism, in which dysglycemia is one axis among other factors 442 that can impact on complications onset/progression and organ dysfunction. Therefore, the 443 metabolic condition of each subject is approached in an integrated way. Also, we differentiate 444 three types of components: etiological factors, mechanisms and millieu. Each encompasses 445 several factors or axis that are projected in separate 2D planes. We postulate that, by deeply 446 profiling a subject for one type of component, we can place him in the corresponding plan. 447 Furthermore, we postulate that we can predict where the individual is in one plan by knowing the 448 others. Ultimately it will allow placing each individual in a last plan where his metabolic state is 449 known. It is natural that there are groups in the data. However, given the possible combinations 450 of affected mechanisms and organs, it is clear that their number is too high for human 451 understanding.

452 This model differs from McCarthy's palette model in two main points: 1) it considers the path to 453 dysmetabolism and not to hyperglycemia; 2) it separates the different etiological factors from the 454 affected mechanisms and from the internal environment to which the person is exposed on 455 different levels (Figure 3). The different planes are thus projected among themselves, giving us 456 the possibility to know one when we fully evaluate the others. This differentiation can be relevant 457 to prioritize the clinical approach to the individual and to delineate distinct integrated therapeutic 458 and preventive strategies to be adopted in the different planes that nonetheless should be validated 459 in clinical studies.

460 Currently, in therapeutic individualization, therapy is first prescribed to hyperglycemia and then 461 adapted according to the individual characteristics of each patient. In contrast, in precision 462 medicine, the therapeutic approach is chosen after assigning the patient to a group that already 463 considers the individual specificities. For example, in the individualized treatment of T2D, a 464 subject without atherosclerotic disease or CKD but with hypertension and poorly controlled 465 glycemia, when on metformin, can be medicated with one of five drugs (DPP4, GLP-1, SGLT-2, 466 thiazolidinediones, sulfonylureas). This will be chosen by each doctor considering some 467 characteristics of the patient, such as weight. In addition, an antihypertensive is associated. In real 468 life, situations are not so clear as in guidelines. For instance, what to do with a patient with T2D 469 on metformin, with good glycemic control (average HbA1c 6.8%) but with evidence of early 470 DKD and without other metabolic risk factors? How intensive and with which agents should he 471 be treated to have the best health outcome? Is it better to use a SGLT-2 inhibitor or/and start an 472 ACE2 inhibitor? Is this the best treatment for all the patients in this condition? Or what to do with 473 another patient with 15 years of T2D, mostly with poor control (HbA1c >8.5%) under different 474 antihyperglycemic medication, without other risk factors or evidence of Diabetes complications? 475 Should we keep trying to put him in a good track of glycemic control? For what purpose? In a 476 precision medicine approach, he would first be assigned into a group of people sharing common 477 features of the overall metabolic condition, already accounting with all his specificities (including 478 millieu, mechanisms and etiological factors) for which the optimal treatment of that group would 479 be already tested, defined and can then be prescribed for that individual.

In order to train and validate this theoretical model, datasets that consider the overall metabolism and deep phenotyping subjects in the distinct proposed planes are needed. Ultimately this model may be implemented in a decision support system that predicts where people are in their overall metabolism. This would assign the individual to a homogeneous group, eventually unravelling his metabolic footprint.

485

486 8. Conclusion

Precision medicine allows tailoring an approach or treatment to different individuals. In other words, a population is stratified into similar groups, considering relevant characteristics to the condition (e.g. T2D). Doing so for each group an appropriate therapeutic approach is defined. Although precision medicine approaches can make use of genetic data, they can also be based on many other types of clinical data. Observed complexity is solved with the help of mathematical algorithms that stratifies individuals into groups by similarity. 493 In the era of omics and digital health, in which we can extract and deal with thousands of features 494 and use them to tailor care to Diabetes, it is not prudent to limit cluster analysis to a few already 495 preestablished common mechanisms. Furthermore, these new strategies allow us to deal with 496 blood glucose levels as a continuum, together with the overall milieu, surpassing the artificial 497 glycemia-based cut-off approach. By fully profiling subjects regarding genomics, environmental 498 factors and time exposition, we will be able to know which mechanism(s) is(are) affected and 499 is(are) responsible for a dysmetabolic condition. This enables the use of drugs in a precise manner 500 and the discovery of new ones. Additionally, prevention of complications, such as cardiovascular 501 events, may be earlier and more effective. The great big challenge will be identifying which 502 features are relevant to consider precise care and gather the data to perform these analyses. In a 503 global village such as our world, we should gather robust clinical data working in a worldwide 504 consortium.

505

506 Conflict of Interest

507 The authors declare that the research was conducted in the absence of any commercial or financial

relationships that could be construed as a potential conflict of interest.

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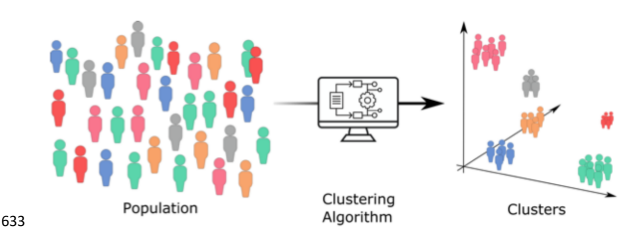
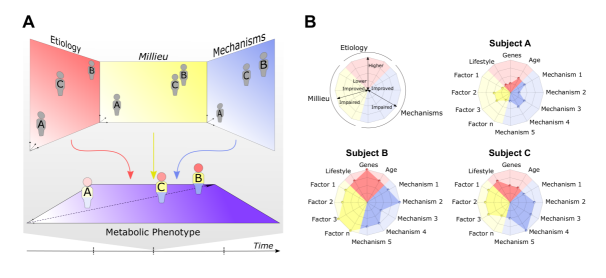


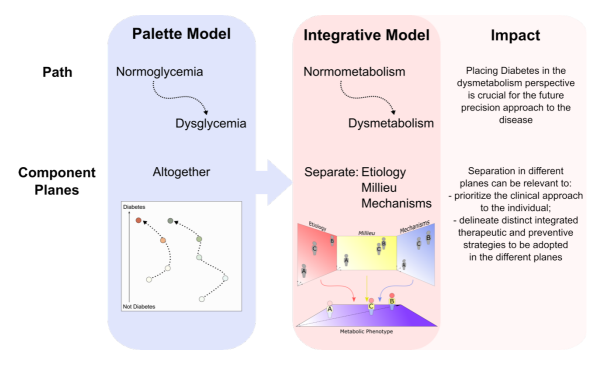
Figure 1 – Cluster analysis scheme. An heterogenous population regarding characteristics of
interest is stratified by a chosen algorithm, that places them in a hyperplane, differentiating natural
homogenous groups.



638

Figure 2 – Integrative model of Diabetes. A) Subjects are deeply characterized regarding etiological factors (including genes, lifestyle and environmental factors), underlying physiopathological mechanisms and metabolic and hemodynamic factors that they are exposed to. They are placed correspondingly onto the Etiology, Mechanisms and Millieu plan. The location of a subject in each plan can be predicted by knowing their position in the others. Ultimately, etiology, mechanisms and milieu projects the subject onto the Metabolic Phenotype

645 plan where its health condition is assessed also considering Diabetes complications as 646 nephropathy, retinopathy as well as cardiovascular complications. Each subject path through time 647 in the Metabolic Phenotype plan can be analyzed but also predicted, leveraging therapeutic and 648 preventive strategies. B) Etiology, Mechanisms and Millieu for each subject can be summarized 649 and more easily visible on a radarplot.



651

Figure 3 – From the Palette Model to the proposed Integrative Model. The Integrative model that
we propose was based on the McCarthys' Palette Model,⁵⁷ but differs essentially in the path and
in the component planes of the model.

655 Tables

Table 1 - Clustering algorithms used in Diabetes studies.

Hierarchical	Partitioning
• Agglomerative ^{26,41}	Hard clustering
	- k-means ⁴¹
	- k-medoids (Partition around Medoids - PAM) ⁵²
	- Self-organising Maps (SOM) 37,41
	Soft Clustering
	- Fuzzy c-mean ⁵⁹

Clustering Algorithm	Advantages	Disadvantages
Hierarchical	 Does not need pre-specification of the number of clusters Accepts any kind of distance function Visualisation of number of clusters Agglomerative good at identifying small clusters, divisive better identifying large clusters 	 High computational cost, it does not scale properly Difficult to alter once the analysis starts Different clusters form according to the linkage function More prone to identify spherical and convex clusters Need to define the cophenetic distance cut-off Sensitive to outliers
k-means	Simple to implement and understandFast and efficient for large datasets	 Require specification of the number of clusters Sensitive to the randomly chosen seeds Some implementations use only More prone to identify spherical and convex clusters
PAM	 Simple to understand and implement Less sensitive to noise and outliers than k-means Allows using general dissimilarities of objects 	 Require specification of number of clusters Sensitive to random initialization of medoids Higher computational cost than k-means More prone to identify spherical and convex clusters Does not scale well for large datasets
SOM	 Easy to understand and interpret Deals with large and complex data sets Finds different clusters formats 	 Many parameters to be set and optimised Computational expensive When initialized randomly, it is sensitive to the initial seeds The number of clusters must be previously defined
b-NMF	 Best results for an overlapped data set Datapoint may belong to more than one cluster. tion around medoids: SOM: Self-Organizing I 	Require specification of the number of clustersComputational cost

658 Table 2 – Advantages and drawbacks of clustering algorithms (Adapted from ²⁶).

659 PAM: partition around medoids; SOM: Self-Organizing Maps.