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1	Serological Prot	eome Analysis F	Reveals New Specific	Biases in the IgN	I and IgG Autoantibody

- 2 Repertoires in Autoimmune Polyendocrine Syndrome Type 1
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4 Running title: Biases in Autoimmune Repertoires in APS 1

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29	Abbreviations:
30	AIRE, autoimmune regulator
31	APECED, autoimmune polyendocrinopathy candidiasis ectodermal dystrophy
32	APS 1, autoimmune polyendocrine syndrome type 1
33	BCR, B cell receptor
34	eTAC, extra-thymic Aire-expressing cells
35	mTEC, medullary thymic epithelial cells
36	OAE, other autoimmune endocrinopathies
37	TS-Ag, tissue-specific antigen
38	
39	Keywords: AIRE; APS 1; autoantibody repertoire; post translational modifications; T-dependent and
40	T-independent mechanisms
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44	

45 Abstract:

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47 **Objective**: Autoimmune polyendocrine syndrome type 1 (APS 1) is caused by mutations in the AIRE 48 gene that induce intrathymicT-cell tolerance breakdown, which results in tissue-specific autoimmune 49 diseases. **Design**: To evaluate the effect of a well-defined T-cell repertoire impairment on humoral 50 self-reactive fingerprints, comparative serum self-IgG and -IgM reactivities were analyzed using both 51 one- and two-dimensional western blotting approaches against a broad spectrum of peripheral tissue 52 antigens. Methods: Autoantibody patterns of APS 1 patients were compared with those of subjects 53 affected by other autoimmune endocrinopathies (OAE) and healthy controls. Results: Using a Chi-54 square test, significant changes in the Ab repertoire were found when intergroup patterns were 55 compared. A singular distortion of both serum self-IgG and self-IgM repertoires was noted in APS 1 56 patients. The molecular characterization of these antigenic targets was conducted using a proteomic 57 approach. In this context, autoantibodies recognized more significantly either tissue-specific antigens, 58 such as pancreatic amylase, pancreatic triacylglycerol lipase and pancreatic regenerating protein 1α , or 59 widely distributed antigens, such as peroxiredoxin-2, heat shock cognate 71-kDa protein and aldose 60 reductase. As expected, a well-defined self-reactive T-cell repertoire impairment, as described in APS 61 1 patients, affected the tissue-specific self-IgG repertoire. Interestingly, discriminant IgM reactivities 62 targeting both tissue-specific and more widely expressed antigens were also specifically observed in 63 APS 1 patients. Using recombinant targets, we observed that post translational modifications of these 64 specific antigens impacted upon their recognition. Conclusions: The data suggest that T-cell-65 dependent but also T-cell-independent mechanisms are involved in the dynamic evolution of 66 autoimmunity in APS 1.

67

68 Introduction

69 Autoimmune-polyendocrinopathy-candidiasis-ectodermal dystrophy (APECED; OMIM ID: 70 601240; 240300), also known as "Autoimmune Polyendocrine Syndrome type 1" (APS 1), is a rare 71 monogenic autosomal recessive disease associated with autoimmune regulator (AIRE) gene mutations 72 (1, 2). The AIRE gene is located on chromosome 21(21q22.3) and encodes the Aire protein, which is 73 expressed in thymic medullary epithelial cells (mTECs), but also in dendritic cells and monocytes (1) 74 in spleen and lymph nodes (3). Aire expresses many structural and functional characteristics common 75 to transcriptional regulators (4). Experimental models using Aire^{-/-} animals have shown that Aire is 76 involved in intrathymicT-cell-negative selection because it promotes ectopic expression of a subset of 77 peripheral tissue-specific antigens (TS-Ags) by mTECs (5, 6). Loss of Aire-dependent thymic 78 expression of a peripheral TS-Ag (such as mucin-6) results in autoimmune reactivity against this 79 protein (7). Aire is also expressed in extra-thymic Aire-expressing cells (eTACs), like myeloid and 80 lymphoid cells (8), in lymph nodes and spleen, where it regulates a set of TS-Ags, suggesting that Aire 81 expression has broad transcriptional consequences for TS-Ag presentation in the periphery. 82 Interestingly, the genes regulated by AIRE in eTACs had no overlap with AIRE-regulated genes in the 83 thymus, suggesting a complementary role in the maintenance of self-tolerance (3). Aire^{-/-} mice 84 develop tissue-specific autoantibodies (autoAbs) and lymphocyte infiltrates in multiple organs (5, 6). 85 In humans, APS 1 is characterized by several tissue-specific autoimmune diseases (9) associated with 86 organ-specific but also non-organ-specific autoAbs detected in the serum (10). Antibodies against 87 cytokines have also been reported (11, 12). APS 1 patients develop endocrine autoimmune diseases, 88 such as adrenal insufficiency, hypoparathyroidism, hypogonadism, type 1 diabetes mellitus and 89 Hashimoto thyroiditis, and non-endocrine autoimmune diseases, such as pernicious anemia, hepatitis, 90 alopecia, vitiligo and candidiasis (9). Thus, APS 1 represents a unique monogenic human model in 91 which a well-demonstrated T-cell tolerance breakdown occurs that can result in several tissue-specific 92 autoimmune diseases. Although numerous APS 1tissue-specific target antigens have been described, 93 the global systemic self-antibody repertoire remains to be defined. We have previously performed 94 such a global immunoproteomic approach in healthy subjects and in patients with different

95 autoimmune diseases (13–15). In healthy subjects, the human Ab repertoire is thought to be well 96 conserved and restricted to a few self-antigens in homologous tissues (16). Nevertheless, in each 97 subject, singular patterns were found, possibly related to individual responses against exoantigens 98 (13). Interestingly, a distortion of serum self-IgG patterns in organ-specific autoimmune diseases that 99 predominantly involve T cells, such as multiple sclerosis, has been demonstrated in our laboratory 100 (13). The intriguing aspect of these data was that discriminant reactivities were supported by widely 101 distributed antigens (14). To define more precisely the pathophysiological significance of these 102 fingerprints, we evaluated, in the present study, the autoreactive antibody response in a pathology 103 where well-defined molecular defects, related to tolerance induction processes, have been described. 104 To evaluate T-cell-dependent and T-cell-independent involvement in APS 1, we compared self 105 immunological patterns obtained with both IgG and IgM autoAbs and characterized the respective 106 molecular targets recognized.

107

108 Subjects and Methods

109 **Patients**

110 Sera from 48 patients were analyzed. The samples were obtained from 14 patients with APS 1 111 (group 1: 9 males, 5 females, mean age = 33 ± 14 years), 17 patients with other autoimmune 112 endocrinopathies (OAE) (group 2: 6 males, 11 females, mean age = 47. 6 ± 15.1 years) and 17 healthy 113 controls (group 3: 9 females, 8 males, mean age = 33.1 ± 9.2 years). APS 1 patients were clinically 114 diagnosed according to Neufeld criteria (17) and confirmed by DNA sequencing as described 115 previously (18). OAE patients presented either single or multiple endocrinopathy (Table 1). Sera were 116 collected with the subjects' written consent and the study was approved by the local ethics committee. 117 118 **Anti-cytokine ELISA** 119 Serum reactivities towards interleukin (IL)-22, IL-17A, IL-17F, IFN-omega and IFN-alpha2A

(all from R and D Systems, Minneapolis, MN), were evaluated in both APS 1 and OAE patients using
ELISAs, as previously reported (12), with either anti-human IgG or IgM alkaline phosphatase-

122 conjugate (Sigma-Aldrich, Poole, UK) as the secondary antibody.

123

124 Tissues

Tissue samples were extracted from post-operative fragments with the patients' written consent or from post-mortem samples. Adrenal tissue was obtained from adrenalectomies performed during nephrectomies for kidney adenocarcinoma; none of the adrenal tissues had been invaded by tumors. Pancreas samples were obtained during postmortem dissections within 6 hours of death and were performed with the approval of the local ethics committee.

130

131 Western blotting and related analytical procedures

One-dimensional electrophoresis (1-DE) or two-dimensional electrophoresis (2-DE) was performed as described in (14). For immunostaining, the gels were blotted onto Hybond-P PVDF membranes (Amersham Pharmacia Biotech Europe GmbH, Saclay, France) using a semidry protocol (8 mA per cm²) as in (14). Dilution of secondary antibodies coupled to peroxidase was1/5000 for antihuman Fc μ and 1/10000 for anti-human Fc γ antibodies. Superimposition and alignment of antibody reactivity was performed using Diversity Database Fingerprint software (version 22; BioRad, Hercules, CA, USA) for 1-DE and PDQuest software (BioRad) for 2-DE.

139

140 **Two-dimensional electrophoresis (2-DE)**

141 Tissue homogenization and 1-DE protein separation were done as previously described (11). 142 Briefly, 100 mg of each tissue was homogenized in a detergent solution (4% Triton X100, 1X anti-143 protease cocktail; Sigma, St Louis, MO, USA) and ground using a grinding kit (GE Healthcare) before 144 protein precipitation with a 2D cleanup kit (GE Healthcare). The supernatant was removed and the 145 pellet was resuspended in 250 ml of sample buffer (8 M urea/2 M thiourea [Sigma], 4% CHAPS 146 [Sigma]). Protein concentration was determined using the Bradford assay (BioRad). Proteins (500 mg 147 per gel) were eluted into rehydration buffer (8 M urea/2 M thiourea [Sigma], 2% CHAPS [Sigma], 148 DeStreak reagent [15 mg/ml, GE Healthcare] and ampholytes [1% IPG buffer, GE Healthcare]) before 149 first separation according to their isoelectric points along a nonlinear immobilized pH-gradient (IPG) 150 strip (pH 3–11 NL, 18 cm long) using an IPGphor III apparatus (GE Healthcare), as described 151 elsewhere (14). For the second dimension, equilibrated strips were loaded onto 8–18% SDS-152 polyacrylamide gels and electrophoresis was performed as in (19). One preparative gel was stained 153 with CBB G-250 (Sigma) and used for spot cutting and protein sequencing. The remaining gels were 154 electroblotted onto ECL membranes (GE Healthcare).

155

156 In-gel digestion and MALDI-TOF/TOF MS analysis

Protein identification was performed using a ProteineerTM workflow from BrukerDaltonics 157 158 (Bremen, Germany). Colloidal Coomassie blue-stained spots were excised from preparative 2D gels 159 using a spot picker (PROTEINEER spTM) and placed onto 96-well microtiter plates. In-gel digestion 160 and sample preparation for MALDI-TOF/TOF analysis were performed according to the 161 manufacturer's instructions using a digester/spotter robot (PROTEINEER dpTM) and a digestion kit 162 (DP 96 standard kit, BrukerDaltonics). The MALDI target plate (AnchorChip[™], BrukerDaltonics) 163 was covered with a cyanohydroxycinnamic acid (CHCA) matrix (0.3 mg/ml in acetone:ethanol, 3:6 164 v/v). Extracted peptides were applied directly onto the CHCA matrix thin layer. The molecular mass 165 measurements were performed in automatic mode using FlexControlTM 22 software on an UltraflexTM 166 TOF/TOF instrument (BrukerDaltonics), in the reflection mode for the MALDI-TOF peptide mass 167 fingerprint (PMF) and in LIFT mode for the MALDI-TOF/TOF peptide fragmentation fingerprint 168 (PFF). External calibration was performed using a peptide calibration standard kit (BrukerDaltonics). 169 Peak lists were generated from MS and MS/MS spectra using FlexanalysisTM 24 software 170 (BrukerDaltonics). Database searches using Mascot (Matrix Science Ltd, London, UK) and PMF 171 datasets were performed via ProteinScape 13 (BrukerDaltonics). Searches were conducted for 172 monoisotopic peptide masses using the NCBI and Swiss-Prot protein databases and Mascot 173 (www.matrix-science.com). Various parameters were used for database searches: mammal species, 174 one missed cleavage, chemical partial modifications (oxidation of methionines, cysteines modified by 175 carbamidomethylation) and a mass tolerance of 75 ppm and 0.5 Da for fragment ions. Criteria used to 176 accept the identifications included the probability score and the number of matched peptides 177 (minimum of 6 peptides).

178

179 **Recombinant proteins**

GST-tagged full length recombinant proteins were purchased from ABNOVA (Aachen, Germany): pancreatic amylase (AMY2A, AAH07060); pancreatic triacylglycerol lipase (PNLIP, AAH14309.1); pancreatic regenerating protein 1 α (REG1A, AAH05350); aldose reductase (AKR1B1AAH00260); peroxiredoxin 2 (PRDX2, AAH00452.1); heat shock cognate 71-kDa protein (HSPA8, AAH16179). They were all produced in wheat germ cell-free system. Five micrograms were loaded on SDS-PAGE mini-gels (Biorad), and were processed as described earlier in the western blotting procedure.

187

188 Statistical analysis

Data were expressed in binary mode (0 = absence of an antigenic band; 1 = presence of an antigenic band) to analyze IgG and IgM antibody patterns using the Chi-square test (a p value "P"< 5.10^{-2} was judged as significant). This approach enabled us to select antigens indicative of qualitatively different immune recognition among the 3 groups, within groups 1 and 2 and within groups 1 and 3.

194

195 **Results**

196 Validation of serum and western blotting procedure

Anti-cytokine antibody reactivity (anti-IFN alpha2A, IFN-omega, anti IFN-lambda1, anti-IL-17A anti-IL-17F, and anti-IL-22) was -evaluated in APS 1 and OAE patients, in order to qualify the sera of the 2 groups (Supplemental Table 1). IgM anti-cytokine reactivity and IgG anti-IL-17A were never observed neither in OAE, nor in APS-1 patients. By contrast, IgG anti-IFN-alpha2A, anti-IFN-0 omega, anti-IL22 and anti-IL-17F were predominantly observed in APS 1 compared to OAE patients (90%, 80%, 70% versus 27%, 18% and 0%, respectively). Anti-IFN-lambda1 Ab (IgG or IgM) were never observed in APS 1 or OAE patients (data not shown).

To evaluate the preservation of relevant antigenic targets after the protein extraction procedure and to test the quality of pancreatic and adrenal tissues as selected targets in this present work, we first 206 evaluated the ability of monoclonal IgG antibodies to detect representative antigenic markers of these 207 tissues. As expected, glutamic acid decarboxylase 65 (GAD 65) and steroid 21 hydroxylase 208 (210Hase) expression was respectively preserved in pancreatic or adrenal tissues. However, NACHT 209 leucine-rich-repeat protein 5 (NALP 5) expression was not observed in these 2 tissues, in contrast to 210 the parathyroid tissue (see Supplemental Figure 1).

211 To enlarge the spectrum of analysis of T-cell-independent and T-cell-dependent self-reactive 212 Abs, the reactivity of the 2 isotypes IgM and IgG were respectively evaluated towards adrenal and 213 pancreatic protein extracts. A similar analysis was preliminarily performed with sera collected in 214 healthy subjects. As illustrated in Figure1A, each isotype was able to recognize protein bands whose 215 expression was shared by the 2 tissues (e.g. ~37 kDa for IgG; ~25 kDa for IgM; black arrows). In 216 addition, each isotype recognized tissue-specific antigens (~50 kDa for IgG in adrenal tissue; ~63-65 217 kDa for IgM in pancreatic tissue; white arrows in Figure1, A and B). Moreover, a same tissue-specific 218 antigenic band was recognized by the 2 isotypes (~20 kDa for adrenal tissue; ~60 kDA for pancreatic 219 tissue; black arrows in Figure1B).

220

221

Serum self-IgG and -IgM reactivities restrictively observed in APS 1 patients

222 As shown in Figure 2, serum self-IgG and -IgM responses against adrenal and/or pancreas 223 protein extracts were quantitatively (numbers of bands) and qualitatively (molecular mass of 224 recognized bands) heterogeneous within subjects, indicating that inter-individual variability occurs. 225 Firstly, there were more antigenic bands in patients with APS 1 or OAE than in healthy controls. 226 When we considered all the sera studied, serum self-IgM reactivity was quantitatively greater than 227 self-IgG reactivity against both extracts, in both groups of patients. For pancreatic extracts, 82 228 different antigenic bands were identified in self-IgM patterns, while 56 bands were noted in self-IgG 229 patterns (P<0. 001). Moreover, for adrenal extracts, 71 antigenic bands were identified in self-IgM 230 patterns, while only 45 bands were noted for self-IgG patterns (P<0. 001). A similar difference was 231 observed in all groups (APS 1, OAE and healthy subjects). In addition, some antigenic bands detected 232 either in adrenal or in pancreatic tissues were common in all sera collected (black arrows in Figure 2, A 233 and B).

234 Marking of antigenic bands related to 210Hase for adrenal extracts and GAD65 for pancreatic 235 extracts revealed a co-alignment of bands only detectable in some APS 1 and OAE patients and never 236 detectable in healthy subjects (see Supplemental Table 2). Immune reactivity against 21OHase or 237 GAD65 was only observed when patients presented adrenal or pancreatic autoimmune diseases. 238 Compared to classical assays performed in routine to define specific Abs towards theses Ag, western 239 blotting procedure is less sensitive. In spite of the singularity found in each pattern, variabilities 240 related to a specific group were observed. Thus, some antigenic bands were only detected on adrenal 241 and/or on pancreatic extracts in APS 1 patients when IgG and/or IgM isotypes were evaluated. We 242 then focused statistical analysis (i) on reactivities observed on adrenal tissue in APS 1 and OAE 243 patients when adrenal insufficiency occurred in these 2 groups, and (ii) on reactivities observed on 244 pancreatic tissue in APS 1 and OAE patients when pancreatic insufficiency occurred in these 2 groups. 245 In these 2 situations, specific reactivities towards either adrenal or pancreatic tissues were specifically 246 observed in APS 1 group.

247 The presence or absence of protein bands of reactivity was evaluated for each tissue. 248 Computer-assisted alignment and additional statistical studies allowed us to localize singular IgG 249 and/or IgM bands of reactivity detected in adrenals (p66, p36 and p25) and in pancreas (p55, p53 and 250 p22) (thin black arrows in Figure 2, A and B). For self-IgG patterns (see Supplemental Figure 2A), the 251 Chi-square test identified 2 bands unique to adrenal tissue (Ad p36 and Ad p25) and 2 bands unique to 252 pancreatic tissue (Pc p55 and Pc p53), which were significantly more often recognized by APS 1 253 patients, compared to OAE patients and healthy controls. For self-IgM patterns (see Supplemental 254 Figure 2B), the Chi-square test identified 2 bands unique to adrenal tissue (Ad p66 and Ad p25) and 2 255 bands unique to pancreatic tissue (Pc p53 and Pc p22), which were significantly more often recognized 256 by APS 1patients compared to the other two groups. Thus two protein bands (Ad p25 and Pc p53) 257 were recognized both by IgG and IgM.

258

259 Characterization of discriminant antigenic bands of reactivity

260 To further characterize the discriminant bands of reactivity, we used a serological proteomic 261 approach. Identification of antigenic targets of such reactivities was first performed by comparing 1-D and 2-D immune patterns. Sera were used to identify antigenic candidates on a proteomic map obtained after 2-D electrophoresis performed for each tissue. Two-dimensional electrophoresis followed by immunoblotting revealed the presence of multiple antigenic spots for pancreatic (Figure 3) and adrenal protein extracts (Figure 4). Then, superimposition of antigenic spots and protein spots revealed by a standard colloidal Coomassie blue-stained two-dimensional gel electrophoresis enabled us to select proteins for further in-gel digestion and MALDI-TOF/TOFanalysis on the basis of peptide mass matching (19).

269 This approach enabled us to identify some proteins as potent discriminant antigens for each 270 tissue using the SWISS-PROT database (Table 2). The antigens identified in our study were either 271 tissue-specific or ubiquitous proteins. Some antigens were targeted both by self-reactive IgM and IgG 272 reactivities: one for adrenal tissue, and 2 for pancreatic tissue. As shown in (Figure 3 and Table 2), 273 pancreatic (Pc) p55 was characterized as pancreatic a-amylase (P04746), Pc p53 as pancreatic 274 triacylglycerol lipase (P16233), and Pc p22 as pancreatic regenerating protein 1α (P05451). 275 Furthermore, adrenal (Ad) p66 was characterized as heat shock cognate 71-kDa protein (P11142), Ad 276 p36 as aldose reductase (P15121) and Ad p25 as peroxiredoxin-2 (P32119) (Figure 4 and Table 2).

277

Evaluation of discriminant reactivities observed in APS 1 patients against targeted recombinant proteins

In order to strengthen the data observed on tissue-extracted proteins, we performed 1-D blotting with recombinant proteins identified by the 2-D approach. Both IgG and IgM reactivities were tested in all APS 1 sera against the 6 discriminant antigens.

For the IgG isotype, and except for lipase, the autoreactivity frequency was similar between tissue protein extracts and recombinant proteins (Table 3). Moreover, in terms of presence or absence of reactivity, concordance levels were higher than 80% when IgG reactivities were evaluated either on tissue extract or recombinant proteins. For the IgM isotype, whatever the frequencies observed, the concordance in terms of presence or absence of reactivity was lower than 75% (Table 3).

288

289 **Discussion**

290 In previous studies, specific serum autoAbs were usually investigated by techniques using 291 purified self-Ags and/or relevant peptides from preselected targets (20). We previously demonstrated 292 the value of using a large panel of antigens derived from different tissue extracts to analyze the serum 293 autoAb repertoire in organ-specific autoimmune disease (13, 15). Using this approach, we illustrated 294 that the specific antibody response associated with a pathological condition could be rich and diverse 295 and not only focalized on a restricted set of antigenic targets. Among the different reactivities, some of 296 them could constitute a specific pathological signature of the disease. In this study, we aimed to 297 analyze the diversity of the autoantibody repertoire in APS 1, not for diagnostic purposes, but to 298 appreciate potential biases specifically associated with this condition. APS 1 is classically described as 299 an Aire-mediated T-dependent disease. In this view, we performed a specific analysis of both the IgG-300 and IgM-specific auto-antibody repertoires in APS 1 patients versus controls, to evaluate the specific 301 distortion restricted to the IgG repertoire in this disease, whereas the IgM patterns were attempted not 302 to be changed.

303 We have first evaluated anti-cytokine Ab reactivity in order to assess whether our patients are 304 representative of both APS 1 and OAE patients reported elsewhere (12). We chose then to analyze the 305 autoAb repertoire on both adrenal and pancreatic protein extracts based on the large diversity of 306 autoAb specificities observed in the sera of APS 1 patients and controls in these tissues. Other tissue 307 protein extracts have been evaluated (gastric, ovarian, testis, liver, thyroid, parathyroid, skin) (not 308 shown) and reveals less global reactivity or more homogeneous autoreactive patterns between groups 309 of patients, as observed previously (10). Moreover, adrenal and pancreatic tissue protein extracts 310 contained specific antigens which have been described as being targeted by autoAb associated with 311 clinical manifestations reported in APS 1 and other autoimmune endocrinopathies, such 210Hase and 312 GAD 65. We have confirmed that, using monoclonal Abs with our western blotting procedure, we 313 could reveal these specificities in these tissues when Abs were present in serum of individuals. As 314 expected, we did not reveal NALP5 expression in these 2 tissues, whereas it was observed in the 315 parathyroid protein extract. Thus, we could not detect anti-NALP5 autoAbs in this study.

316 We first evaluated the global richness of the immune repertoire. With regard to the IgM 317 autoAb panels, we observed in all individuals a more diversified repertoire than with IgG, which could 318 be related to the natural immune repertoire ('immunculus') that has been described as being largely 319 composed of IgM autoAbs (21). By contrast, we observed a more diversified IgG autoAb repertoire in 320 patients suffering from autoimmune diseases, namely APS 1 and OAE, than in healthy subjects. This 321 phenomenon could be related to at least two events. First, a specific defect of the educational process 322 of the immune system leading to the persistence of autoreactive immune cells could contribute to the 323 enlargement of this autoreactiveAb repertoire in patients. Secondly, the tissue damage associated with 324 the autoimmune process could contribute to enlarging the panel of autoantigens that are expressed in 325 altered tissues and presented to these immune cells. This phenomenon may itself contribute both to the 326 preservation of autoimmune specificities and to the emergence of new autoreactiveAb specificities, 327 generating a neo-repertoire. These two processes may also act by a summation effect.

328 In a second step, we analyzed the intra-individual variations of the autoreactive patterns 329 between adrenal and pancreatic tissues. We observed that some IgM or IgG reactivities were co-330 aligned between the two tissues, suggesting that a cluster of widely distributed auto-antigens could be 331 targeted by these reactivities. By contrast, some bands of reactivity were exclusively observed on 332 either the adrenal or the pancreatic protein extracts, suggesting tissue-specific autoimmune targeting. 333 We next studied the inter-individual variations of the autoreactive patterns independently on the 334 adrenal or pancreatic tissues. Whereas we did not observe any difference in terms of number of bands 335 of reactivity between APS 1 and OAE, we hypothesized that qualitative distortions could be 336 specifically associated with the Aire-related pathological process in APS 1. This condition is described 337 as a T-dependent autoimmune disorder which preferentially impacts the IgG autoAb repertoire. 338 Surprisingly, in APS 1 patients compared to the 2 control groups, our approach demonstrated as much 339 as specific distortions in IgM repertoire that in IgG repertoire. In our study, some IgM specificities 340 were shared by different APS 1 patients, suggesting that the autoreactivities supported by this isotype 341 are sustainable and not transitory reactivity brought to switch to the IgG class. Several studies have 342 reported an extrathymic expression of Aire that influenced the T-cell repertoire(3, 5). Our data suggest 343 that Aire expression deficiency in peripheral lymphoid organs could also impact the autoreactive IgM 344 repertoire. Sustained IgM production by B cells has been associated with two different B-cell 345 subpopulations in humans. It has been reported that during germinal center differentiation, follicular B cells could maturate into long-lasting IgM-expressing memory B cells through T-dependent mechanisms (22). By contrast, T-independent mechanisms generate marginal zone B cells which produce IgM in response to non-peptidic epitopes (23, 24). Interestingly, numerous studies have focused on changes affecting B-cell homeostasis and T-cell-independent marginal zone (MZ) B-cell subsets in Aire^{-/-} mice (25–28). In addition, recent studies have underlined the fact that Aire can regulate T-cell-independent B-cell responses through B-cell-activating factor of the TNF family (BAFF) (28).

353 Using a serological proteomic approach, we did not observe any discriminant reactivities 354 towards 210Hase or GAD specifically associated with APS 1 condition. These results could be related 355 to the presence of common reactivities in the control group of patients with other autoimmune 356 endocrinopathies, since both antibodies could be observed in both APS 1 and other 357 polyendocrinopathies. By contrast, we noted that some reactivity were statistically more observed in 358 the APS 1 group. They targeted tissue-specific antigens such as amylase, lipase and pancreatic 359 regenerating protein 1 alpha. They also recognized three ubiquitous antigens: peroxyredoxine-2, heat 360 shock cognate 71-kDa protein and aldose reductase. The discriminant recognition of amylase, lipase 361 and pancreatic regenerating protein 1 alpha emphasizes pancreatic exocrine dysfunctions widely 362 evoked either in APS 1 patients who could develop malabsorption caused by several mechanisms such 363 as exocrine pancreatic insufficiency (29-35) or in experimental models such as NOD Aire-deficient 364 mice (5, 36).

365 The discriminant targeting of aldose reductase, a ubiquitous protein mainly expressed in 366 adrenal glands (37), by IgG Abs in APS 1 patients, requires consideration. It has been shown that the 367 expression of aldose reductase is regulated by Aire in mTECs in mice (5). In addition, the Aire-368 dependent expression of other ubiquitous Ags has also been described in eTACs localized in lymph 369 nodes and the spleen (3). Multi-organ inflammation in Aire-deficient models is also known to be 370 associated with the presence of serum autoAbs against proteins specifically produced by these organs. 371 In our study, the discriminant targeting of some ubiquitous antigens, such as peroxyredoxine-2 and 372 heat shock cognate 71-kDa protein, could be indicative of endogenous danger signals involving 373 cellular oxidative stress. It can be compared to biomarkers previously described in systemic autoimmune disorders (38, 39).

375 In parallel with the combination of reactivities classically associated with APS 1 diagnosis, 376 our data highlight some biomarkers that could be associated with a particular tissue alteration 377 (exocrine pancreatic-specific antigens) or more general pathological processes associated with 378 autoimmune diseases. We aimed to design an in vitro assay to evaluate the presence of these 379 reactivities, using recombinant antigens. IgG reactivities towards amylase, aldose reductase and 380 peroxyredoxine-2 were also observed in APS 1 patients when we used recombinant proteins as targets. 381 By contrast, IgM reactivities against HSP71, REG-1A, and lipase were not concordantly observed 382 between tissue extracts and recombinant proteins. When detected, such IgM reactivities were 383 systematically observed at a lower frequency when we used recombinant proteins. Such a discrepancy 384 between the immunoproteomic approach and an in vitro assay using recombinant targets has 385 previously been observed (40). To avoid the impact of folding on antigenic recognition, we chose to 386 use the same one-dimensional electrophoresis experimental procedure. The denaturing conditions lead 387 to the linearization of proteins whatever their origin: tissue extracts or purified wheat germ 388 recombinant proteins. Nevertheless, the presence of post transcriptional modifications (PTMs), such as 389 glycosylation, on the targeted antigens could support these observations. Eukaryotic PTMs are not 390 observed in the wheat germ expression system, so that specific modifications of native proteins are not 391 present on the recombinant protein. Interestingly, we observed a major reactivity discrepancy when we 392 focused on the IgM isotype. Once again, these observations could highlight the impact of the thymo-393 independent processes associated with the dynamic changes in the IgM repertoire in APS 1 patients.

394 Conventional immunoassays are usually performed with limited antigenic targets, the choice 395 of which has been driven by a supposedly well-known physiopathogenic rationale. Advances in 396 proteomic methodologies (in vitro gene expression, 2-DE and mass spectrometry) have allowed the 397 emergence of broad spectrum analysis methods. These approaches have been developed to overcome 398 the limits of conventional methods. Based on a "without any a priori" strategy, they offer a 399 simultaneous analysis of a large spectrum of reactivities, which surpasses the physiopathogenic 400 hypotheses and offers an integrative interpretation of results. When applied to the APS 1 condition, 401 this immunoproteomic methodology not only reveals the expected IgG repertoire biases, it also 402 identifies IgM repertoire distortions. The latter alterations could be partially associated with T-403 independent immunological events related to the impact of post-translational modifications of 404 antigens. Our results highlight the fact that AIRE also impacts the presentation of thymo-independent 405 antigens. It points out that autoimmune alterations observed in APS 1 are not only related to Aire-406 driven T-cell clonal deletion deficiency. At an individual level, this approach highlighted original 407 antigenic targets, potentially associated with tissue injury and cellular dysfunctions related to the 408 singular clinical evolution in each patient.

409

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426	Author contributions
427	Conceived and designed the experiments: SD, EP, DL, LP. Performed the experiments: EP,
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- 551

Patient	Age	AIRE gene mutations	Clinical manifestations					ntibodie			
	(Gender)	AIXL gene initiations	Chinear mannestations	TPO	TG	TBII	GAD	IA2	21 OHase	GPC	tTg
APS 1											
1	37 (F)	c.967_979del13 / c.967_979del13	HPT, AI, DM, OI, PA, C, A, K				+	+	+	+	
2	28 (M)	c.1193delC / c.1193delC	HPT, AI, PA, C, M	+	+				+		
3*	42 (F)	c. 1097 C>T / c. 769 C>T	HPT, AI, OI, PA, C								
4	52 (M)	c.769C>T/c.967_979del13	HPT, AI, C, A, K	+							
5**	31 (F)	c.967_979del13 / c.967_979del13	HPT, AI, OI, PA, C, A						+		
6**	26 (M)	c.967_979del13/ c.967_979del13	С, А, К								
7	25 (M)	c.966_978del13 / c.967_979del13	HPT, AI, C, A						+		
8	23 (M)	c.967_979del13 / c.967_979del13	AI, DM, C, A,M				+				
9	51 (M)	c.769C>T/c.14-1-28G>C	HPT, AI, M, C				+		+		
10	15 (M)	c.967_979del13 / c.967_979del13	AI, C, A, M				+				
11	9 (M)	c.769C>T/c.967_979del13	AI, C, A						+		
12	32 (M)	c.967_979del13/ c.967_979del13	HPT, AI, T, A, K, C	+	+		+		+		
13	39 (F)	c.967_979del13 / c.967_979del13	AI, OI, C		+						
14*	57 (F)	c. 1097 C>T / c. 769 C>T	HPT		+						
OAE			-								
15	25 (F)	ND	T, OI	+	+						
16	25 (F)	ND	DM, PA, M				+	+			+
17	60 (M)	ND	HPT, T, PA	+	+					+	
18	62 (F)	ND	T, OI, PA	+	+					+	
19	57 (F)	ND	DM, T, PA	+	+		+			+	
20	74 (F)	ND	AI, DM, T	+	+				+		
21	56 (F)	ND	DM, T, M	+	+		+	+			+
22	23 (F)	ND	Т	+							
23	63 (M)	ND	AI, DM, T	+	+				+		
24	47 (F)	ND	AI, T	+	+	+			+		
25	43 (F)	ND	AI, DM, T	+	+	+					
26	54 (F)	ND	AI, T	+	+				+		
27	48 (M)	ND	DM, M								
28	35 (F)	ND	Т	+	+	+					
29	61 (M)	ND	T, PA	+	+	+				+	
30***	39 (M)	ND	AI						+		
31***	37 (M)	ND	AI								

 Table 1.Main Clinical Characteristics of APS 1 and OAE patients.
 AIRE gene mutations are detailed in APS 1 patients.

Clinical manifestations: HPT: Hypoparathyroidism; AI: Adrenal insufficiency; DM: Diabetes mellitus; T: Thyroiditis; OI: Ovarian insufficiency; PA: Pernicious anemia; M: Malabsorption; K : Keratitis; A: Alopecia; C: Candidiasis.

Specific Antibodies: TPO: anti-thyroperoxidase; TG: anti-thyroglobulin; TBII: thyroid-binding inhibitory immunoglobulin; GAD : anti-glutamic acid decarboxylase 65

(GAD 65); IA2: antityrosine phosphatase; 21 OHase : anti-steroid 21 hydroxylase ; GPC: anti-gastric parietal cells; tTg: anti-tissular Transglutaminase

M = male, F = female; ND = not done; * siblings, ** siblings, *** siblings.

Name of antigenic band	Isotype concerned	UniProtKB/ Swiss-Prot references	Name	Theoretical mass (Observed mass) (kDa)	Theoretical IP (Observed IP)	MS Mascot Score	MS Sequence coverage	MS/MS Mascot Score	MS/MS Sequence coverage
ADRENA	L TISSUE								
Ad p66	IgM	P11142	Heat shock cognate 71-kDa protein	70.7 (66)	5.4 (5.3-5.5)	169	33.3	217;8	6.32
Ad p36	IgG	P15121	Aldose reductase	35.7 (36)	6.6 (6.1-6.9)	191	50.2	132.7	12.69
Ad p25	IgM/IgG	P32119	Peroxiredoxin-2	21.7 (25)	5.6 (5.6)	190	53.3	537.8	33.5
PANCREAT	TIC TISSUE								
Pc p55	IgG	P04746	Pancreatic α-amylase	57.7 (55)	6.7 (7.1)	201	45.6	338	2.77
Pc p53	IgM/IgG	P16233	Pancreatic triacylglycerol lipase	49.5 (53)	6.2 (7.0)	187	68.9	561.7	16.7
Pc p22	IgM	P05451	Pancreatic regenerating protein 1 alpha	18.7 (22)	5.6 (5.5)	120	54	365.5	30.7

Table 2. Characterization of Discriminant Antigens by MS and MS/MS. Antigens Preferentially Recognized by APS 1 Patients (Chi-2 test results).

		Autor	eactivity frequency	Concordance level		
Target	Isotype	in tissue	on recombinant protein	Presence of reactivity	Absence of reactivity	
Adrenal tissue						
Heat shock cognate 71-kDa protein	IgM	50%	50%	60%	40%	
Aldose reductase	IgG	40%	50%	100%	85%	
Peroxiredoxine-2	IgG	50%	60%	100%	80%	
Peroxiredoxine-2	IgM	40%	40%	75%	85%	
Pancreatic tissue						
Pancreatic alpha-amylase	IgG	100%	90%	90%	-	
Deperantia trigovialvagral lingas	IgG	100%	40%	40%	-	
Pancreatic triacylglycerol lipase	IgM	50%	30%	30%	71%	
Pancreatic regenerating protein 1 alpha	IgM	50%	0%	0%	100%	

Table 3. Level of Concordance of Autoreactivity According to the Origin of the Antigens (Tissue or Recombinant) and to the Isotype (IgG or IgM)

Supplemental table 1. Frequency of detection of IgG and IgM anti-IFN-alpha2A, anti-IFN –omega, anti-IL-22, anti-IL-17A and anti-IL-17F antibodies in APS 1 and OAE patients

	IFN-al	pha2A	IFN –	omega	IL-	-22	IL-	17A	IL-	17F
	IgG	IgM	IgG	IgM	IgG	IgM	IgG	IgM	IgG	IgM
APS 1 patients	90%	0%	40%	0%	80%	0%	0%	0%	70%	0%
OAE patients	27%	0%	0%	0%	18%	0%	0%	0%	0%	0%

Supplemental table 2. Frequency of detection of a co-alignment with anti 21 OHase and anti GAD65 IgG monoclonal antibodies in APS 1 patients, OAE patients and healthy controls.

	GAD65		210Hase		
	IgG	IgM	IgG	IgM	
APS 1 patients	33%	25%	60%	0%	
OAE patients	40%	26%	46%	0%	
Healthy controls	0%	0%	0%	0%	