Prevalence and phenotype of subjects carrying rare variants in the Italian registry for alpha1-antitrypsin deficiency

I Ferrarotti, J Baccheschi, M Zorzetto, C Tinelli, L Corda, B Balbi, I Campo, E Pozzi, G Faa, P Coni, G Massi, G Stella, M Luisetti

Alfa1-antitrypsin deficiency (AATD) is a genetic condition associated with an increased risk of developing chronic obstructive pulmonary disease (COPD) early in life and, to a lesser extent, liver disease. Significant advances have been made during the last decades in understanding its epidemiology and it has been recently suggested that AATD is one of the commonest inherited disorders not only in Caucasians but also among other ethnic groups worldwide.

Although AAT is a highly pleomorphic glycoprotein, with approximately 100 variants having been identified, two major deficient variants, namely Z and S, account for most cases of AATD, since the vast majority of such individuals carry the PI*ZZ or PI*SZ genotype, coding for approximately 15% and 25%, respectively, of normal AAT plasma levels. The establishment of international registries, including large series of AATD individuals, has allowed not only better definition of the epidemiology of AATD, but also more precise definition of the associated clinical phenotypes.

Nevertheless, there are at least 30 AAT alleles other than the PI*Z and the PI*S alleles which are associated with significantly reduced or absent plasma AAT levels. Given the extreme rarity of such variants, often described in the literature as single case reports, little is known about their epidemiology and even less is known about the associated clinical phenotypes.

The Italian Registry for Severe AATD was established in 1996 as a result of a nationwide screening programme sponsored by the two major Italian scientific respiratory societies. Although Italy is considered a country with a medium-low prevalence of AATD (mean PI*Z gene frequency: 0.0013), the programme succeeded in identifying a relatively large cohort of AATD individuals. During the development of the screening program, we noticed that, in addition to the groups of AATD individuals carrying the PI*ZZ and PI*SZ genotypes, there was an unexpectedly large group of subjects carrying at least one rare AATD allele. We therefore decided to study this group of subjects, focusing particularly on characterising their clinical phenotypes.

METHODS

Screening programme

The targeted screening programme, based on dried blood spots, has already been described in detail. Briefly, paper filters and questionnaires were distributed to respiratory physicians throughout the country. Recommendations for AATD screening were the presence of the following: early-onset COPD, familial clustering of COPD, reduced levels of z1-globulins on electrophoresis, serum levels of AAT <80 mg/dl (nephelometry) or <150 mg/dl (immunodiffusion), or a family history of AATD. Paper filters containing the blood spots were shipped to the Central Phenotyping Laboratory in Rome and submitted to isoelectric focusing. In the case of an abnormal isoelectric focusing pattern, the referring physician was asked to ship a serum sample and a frozen whole blood sample. If the abnormal isoelectric focusing pattern was confirmed, the sample was then investigated at a molecular level. The subject’s demographic and clinical data were retrieved from questionnaires filled in by the referring physicians and shipped together with the specimens. It was recommended that the diagnosis of COPD follow international guidelines. All subjects gave their consent to undergo the genetic investigation, which was approved by the ethical committees of the institutions involved.

Key points

- Most subjects affected by z1-antitrypsin deficiency (AATD) carry the PI*ZZ or PI*SZ genotype. Nevertheless, there are at least 30 AAT alleles other than PI*Z or PI*S associated with reduced or absent plasma AAT levels. Little is known about their epidemiology or associated clinical phenotypes.
- Over 98 months, 2922 subjects enrolled by the Italian Registry for AATD, which conducts a screening programme based on dried blood spots, were screened. A total of 155 subjects with severe AATD were identified (132 index cases), together with 152 individuals with intermediate AATD (84 cases).
- Among subjects with severe AATD, we recorded an 11% prevalence of deficient genotypes other than the common PI*ZZ or PI*SZ (15 out of 132 deficient index subjects identified). Among subjects with intermediate AATD, we recorded a 15% prevalence of deficient genotypes other than the common PI*MZ (13 out of the 84 index cases).
- As the cohort of subjects carrying rare AATD variants was relatively large, we were able investigate their characteristics in terms of associated clinical phenotypes, pulmonary lung function, smoking habit, and geographic distribution. We found that these rare variants had a special position within the spectrum of genotype-phenotype correlations in z1-antitrypsin deficiency.
- The prevalence of rare z1-antitrypsin variants found in Italy is the highest so far recorded worldwide and, interestingly, occurs in a country with a medium-low prevalence of the common PI*ZZ genotype.

Abbreviations: AATD, alpha1-antitrypsin deficiency; COPD, chronic obstructive pulmonary disease; OR, odds ratio; 95% CI, 95% confidence interval
Detection of AATD variants

Genomic DNA was extracted from whole blood cells using the standard technique. The S and Z variants were genotyped by a commercially available amplification-reverse hybridisation test kit (Symbiosis, Coccinato, Italy) and by PCR-RFLP using TaqI as the restriction enzyme. The genomic DNA was sequenced after PCR amplification of all coding exons (II-V). All sequencing products were obtained using the ABI Prism BigDye Terminator Cycle Sequencing Ready Reaction kit (Applied Biosystems, Arrington, UK) and were analysed in an automatic ABI 377 DNA sequencer.

RESULTS

During the 98 months from February 1996 to April 2004, 2922 subjects were screened, thanks to more than 250 physicians throughout the country who shipped at least one paper filter. A total of 155 individuals (5.3%) with severe AATD were detected (table 1). Of these, 132 were index cases. The R/D (rare/deficient) subset identifies subjects with severe AATD (homozygous for the rare AAT variant or compound heterozygous with the common Z deficient variant). The M/R (M/rare) subset consists of subjects with intermediate AATD (one rare AAT variant in combination with the normal M allele). Non-index individuals were identified during family screening.

### Table 1

<table>
<thead>
<tr>
<th>Severe AATD</th>
<th>Intermediate AATD</th>
</tr>
</thead>
<tbody>
<tr>
<td>PI*ZZ (n = 114)</td>
<td>PI*SZ (n = 25)</td>
</tr>
<tr>
<td>Index cases, n (%)</td>
<td>96 (84)</td>
</tr>
<tr>
<td>Non-index cases, n</td>
<td>18</td>
</tr>
</tbody>
</table>

The R/D (rare/deficient) subset identifies subjects with severe AATD (homozygous for the rare AAT variant or compound heterozygous with the common Z deficient variant). The M/R (M/rare) subset consists of subjects with intermediate AATD (one rare AAT variant in combination with the normal M allele). Non-index individuals were identified during family screening.

The majority of the subjects carried the Mmalton and Mprocida variants. Four subjects carried the Plowell variant, two subjects carried the I variant, and one subject each the Mheerlen, Q0procida, Q0cairo, and Q0clayton Variants.

Demographic details are given in table 3. From now on, we will present data from index cases only. Comparisons were performed among individuals homozygous or compound heterozygous for deficiency alleles (that is, R/D, PI*ZZ, and PI*SZ) and between individuals heterozygous for a deficiency allele and a normal allele (M/R and PI*MZ). There were no differences as far as mean age at diagnosis was concerned. Male subjects were more frequent in the R/D group than in the PI*SZ group. Former smokers were significantly more frequent among R/D individuals than among both PI*ZZ and PI*SZ subjects (p = 0.021 and p = 0.013, respectively). Whole cigarette smoke exposure (current+former) was detected in 100% of R/D individuals v 67% of PI*ZZ subjects and 52% of PI*SZ subjects (p = 0.008 and p = 0.0017, respectively), as well as in 85% of M/R subjects v 52% of PI*MZ subjects (p = 0.03). As far as the plasma AAT levels were concerned, the R/D subset had a lower mean (SD) level of 29 (18) mg/dl, similar to that of the PI*ZZ group (28 (11) mg/dl), but significantly lower than that of the PI*SZ group (62 (16) mg/dl; p < 0.0001). The M/R subset had a mean (SD) AAT level of 61 (20) mg/dl, which is significantly lower than that of the related PI*MZ group (93 (23) mg/dl; p < 0.0001).

Tables 4 and 5 present the patients’ phenotypes, that is, the associated clinical manifestations and lung function data, retrieved from questionnaires filled in by the referring physicians.

### Table 2

<table>
<thead>
<tr>
<th>R/D subset</th>
<th>M/R subset</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genotype</td>
<td>Number of subjects</td>
</tr>
<tr>
<td>Mmalton/Mmalton</td>
<td>5 (5)</td>
</tr>
<tr>
<td>Z/malton</td>
<td>3 (3)</td>
</tr>
<tr>
<td>Mprocida/Mprocida</td>
<td>2 (2)</td>
</tr>
<tr>
<td>Z/I</td>
<td>2 (2)</td>
</tr>
<tr>
<td>Z/plowell</td>
<td>2 (1)</td>
</tr>
<tr>
<td>Z/Mprocida</td>
<td>1 (1)</td>
</tr>
<tr>
<td>Z/Q0procida</td>
<td>1 (1)</td>
</tr>
</tbody>
</table>

Index cases are in brackets.
Among subjects with severe AATD, COPD (alone or in combination with chronic liver disease) was diagnosed more frequently in the R/D group (100%), than in the two other related groups, PI∗ZZ (97%), and PI∗SZ (88%, p = 0.004). In seven cases out of 96 PI∗ZZ (7%) and in two out of 15 R/D (13%), COPD and liver disease were diagnosed simultaneously. Prevalence of chronic liver disease did not differ among subjects carrying severe AATD; it was absent in M/R subjects, but it was detected in 11% of PI∗MZ individuals. A number of other associated conditions were reported: with the exception of one PI∗MZ subject diagnosed with Wegener’s granulomatosis, which has been repeatedly associated with AATD, the other conditions are most likely to be chance associations. The percentage of healthy subjects increased significantly among the two groups with intermediate AATD (42% in the PI∗MZ group, and 46% in the M/R group).

With reference to the functional phenotype (table 5), the mean baseline FEV1 value (% predicted) was significantly lower in the R/D group as a whole than in the related PI∗SZ group (p = 0.00095), but not significantly lower than in the PI∗ZZ group. The mean FEV1/FVC among the five genotype groups was consistently different (table 5). However, when individuals with normal lung function (healthy, liver disease without COPD, other conditions) were disaggregated from the whole group, the FEV1 and FEV1/FVC in the remaining patients with COPD no longer differed among the R/D, PI∗ZZ, and PI∗SZ subgroups. The difference between PI∗MZ and M/R subjects for FEV1 was significant, but it may have been influenced by the low number of M/R subjects with COPD.

The effect of smoking habit and genotype on FEV1/FVC is summarised in table 6. Logistic regression and multivariate analyses showed that smoke and PI∗MZ increased significantly among the two groups with intermediate AATD (13%), COPD and liver disease were diagnosed simultaneously. Prevalence of chronic liver disease did not differ among subjects carrying severe AATD; it was absent in M/R subjects, but it was detected in 11% of PI∗MZ individuals. A number of other associated conditions were reported: with the exception of one PI∗MZ subject diagnosed with Wegener’s granulomatosis, which has been repeatedly associated with AATD, the other conditions are most likely to be chance associations. The percentage of healthy subjects increased significantly among the two groups with intermediate AATD (42% in the PI∗MZ group, and 46% in the M/R group).

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The effect of smoking habit and genotype on FEV1/FVC is summarised in table 6. Logistic regression and multivariate analyses showed that smoke and PI∗ZZ and PI∗RD genotypes act as independent risk factors for an FEV1/FVC<0.70.

DISCUSSION

Our finding of a 11% prevalence of subjects with severe AATD carrying genotypes other than PI∗ZZ and PI∗SZ is, to our knowledge, the highest reported so far. The NHLBI Registry for severe AATD with 1021 subjects includes 1.7% with genotypes other than PI∗ZZ or PI∗SZ. The Alpha One Foundation Research Network Registry also includes subjects with intermediate AATD: individuals with rare AATD variants accounted for 5.7% of the total. However, in our series including subjects with both severe and intermediate AATD increased the prevalence to 13%. Thus, our large series of rare AATD variants, detected within the relatively small sized Italian registry, prompted us to investigate the characteristics of these subjects more closely.

Molecular characterisation showed that the majority of subjects with rare AATD variants, including both index and non-index cases, carried at least one Mmalton allele (16/37 individuals, 21/74 alleles). This mutation is raised on the M2 base allele and consists of the deletion of an entire TTC codon in exon II, and subsequent deletion of the Phe51 or Phe52 residue of the mature protein. Ten of the 44 subjects carried at least one Mprocida allele, based on M1(Val213), a T→C point mutation at codon 41 exon II, leading to a proline for leucine substitution. The I allele, found in two individuals, and the Plowell allele (also referred to as QOCardiff), found in four individuals, are both raised on the M1(Val213) base allele. The I allele is characterised by a C→T point mutation at codon 39 exon II, leading to a cysteine for arginine substitution, whereas the Plowell allele is characterised by an A→T transversion at codon 256 exon III, leading to a valine for asparagine substitution. The Mlowen variant, found in one subject, is raised on the M1(Ala213) base allele and it is characterised by a C→T point mutation at codon 369 exon V, leading to a leucine for proline substitution. The QOprocida allele (also referred to as Nullprocida or Nullisola di procida) is a null variant characterised by a 17 kb deletion encompassing exons II–V. The QOlayouto a null variant raised on the M1(Val213) base allele, more recently identified, is characterised by a C insertion in exon V, causing a 3’ frameshift mutation, in turn resulting in a stop codon at residue 376. Finally, two AATD variants were recently found in our series of subjects: the Mvarallo allele was first described in a family from a Northern Italian village.

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**Table 3** Characteristics of the 132 index subjects with severe AATD (PI∗ZZ, PI∗SZ, and PI∗RD) and the 84 index subjects with intermediate AATD (PI∗MZ and PI∗MR)

<table>
<thead>
<tr>
<th>Genotype</th>
<th>COPD, n (%)</th>
<th>Liver disease, n (%)</th>
<th>COPD-liver disease, n (%)</th>
<th>Other conditions, n (%)†</th>
<th>Healthy, n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PI∗ZZ (n = 96)</td>
<td>69 (72)</td>
<td>6 (6)</td>
<td>7 (7)</td>
<td>4 (4)</td>
<td>10 (11)</td>
</tr>
<tr>
<td>PI∗SZ (n = 21)</td>
<td>8 (38)</td>
<td>4 (19)</td>
<td>0</td>
<td>2 (10)</td>
<td>7 (33)</td>
</tr>
<tr>
<td>R/D (n = 15)</td>
<td>13 (87)</td>
<td>0</td>
<td>2 (13)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>P</td>
<td>M</td>
<td>Z (n = 71)</td>
<td>27 (38)</td>
<td>8 (11)</td>
<td>0</td>
</tr>
<tr>
<td>M/R (n = 13)</td>
<td>6 (46)</td>
<td>0</td>
<td>0</td>
<td>1 (8)</td>
<td>6 (46)</td>
</tr>
</tbody>
</table>

*p = 0.026 v PI∗ZZ; **p = 0.012 v PI∗ZZ and p = 0.013 v PI∗SZ; ***p = 0.04 v PI∗MZ.

†Plasma AAT levels are those reported by the referring physician. The large variability is due to the fact that both nephelometric and radial immunodiffusion methods were used; values obtained by the latter method are higher than those obtained by nephelometry.

**Table 4** Characteristics of the clinical phenotype

<table>
<thead>
<tr>
<th>Genotype</th>
<th>COPD, n (%)</th>
<th>Liver disease, n (%)</th>
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<td>0</td>
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</tr>
<tr>
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<td>M</td>
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<td>0</td>
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<td>6 (46)</td>
<td>0</td>
<td>0</td>
<td>1 (8)</td>
<td>6 (46)</td>
</tr>
</tbody>
</table>

*p = 0.0036 v PI∗SZ.

†Category includes: psoriasis, chronic sinusitis, pulmonary fibrosis, coronary artery disease, Wilson’s disease, Wegener’s granulomatosis, idiopathic dilated cardiomyopathy, and pulmonary alveolar proteinosis.
This allele is characterised by a 30 bp deletion accompanied by a 22 bp fragment insertion at the 41–51 codon region in exon II, whereas the Q0cairo is a null variant raised on the by a 22 bp fragment insertion at the 41–51 codon region in

In addition to revealing the high prevalence of rare AAT variants in Italy, which will be discussed later, our relatively large series prompted us to examine the phenotypic profile of carriers of such variants. Taken as a whole and looking at index cases only, there was a higher proportion of men (12/15) and a higher proportion of current or former smokers (100%) among carriers of rare variants than in the other groups with severe AATD (PI*ZZ and PI*SZ) (table 3). Among subjects with intermediate AATD, M/R subjects smoked more than the related PI*MZ subjects (table 3). The higher proportion of former smokers among R/D individuals is intriguing: perhaps they stopped smoking because of the lung disease.

Analysis of COPD prevalence and respiratory function data among the groups revealed interesting findings. We found an increasing prevalence of COPD according to the hierarchy PI*MZ (38%) and PI*SZ (38%)<M/R (46%)<PI*ZZ (79%)<R/D (100%) (table 4). Accordingly, the magnitude of FEV1/FVC ratio impairment followed the same hierarchy: PI*MZ (0.74)<PI*SZ (0.71)<M/R (0.67)<PI*ZZ (0.57)<R/D (0.39) (table 5).

Thus, within the spectrum of prevalence of lung disease associated with AATD, carrying a rare AAT variant(s) places R/D individuals at the highest risk, in the same position as PI*ZZ individuals, whereas M/R individuals are in an intermediate position, with PI*SZ and PI*MZ individuals at the end with lower risk. Since it is widely accepted that phenotypes result from the interaction between genetic determinants (in this case, the plasma levels of AAT) and environmental determinants (in this case, smoking), then the phenotypic ranking seems to result from the interaction between the AAT plasma levels reported in our series, with the R/D and PI*ZZ groups at the lowest end (29 and 28 mg/dl, respectively), followed by PI*SZ (62 mg/dl) and M/R (61 mg/dl), and with PI*MZ individuals at the highest end (93 mg/dl), and the smoking prevalence, ranking R/D (100%)>M/R (85%)>PI*ZZ (67%)>PI*SZ (52%) and PI*MZ (52%). These findings further support the concept that the genetic risk factor for COPD is significantly related to the AAT level, and that cigarette smoking may influence the risk rate in individuals with AATD.

The second most common feature associated with AATD, was detected in 13% of R/D subjects, 13% of PI*ZZ subjects, 19% of PI*SZ subjects, and 11% of PI*MZ subjects (table 4). Diagnosis of chronic liver disease, which is the second most common feature associated with AATD, was detected in 13% of R/D subjects, 13% of PI*ZZ subjects, 19% of PI*SZ subjects, and 11% of PI*MZ subjects (table 4). Diagnosis of chronic liver disease, which is the second most common feature associated with AATD, was detected in 13% of R/D subjects, 13% of PI*ZZ subjects, 19% of PI*SZ subjects, and 11% of PI*MZ subjects (table 4).
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Competing interests: none declared

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REFERENCES


5 Alpha-1-Antitrypsin Deficiency Registry Study Group. Survival and FEV1 decline in individuals with severe deficiency of α1-antitrypsin. Am J Respir Crit Care Med 1998; 158: 49–54


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Letter to JMG

286
25 Coni P, Pili E, Convertino G, Pichiri G, Balestino A, Del Mastro M, Donner CF, 
26 Seersholm N, Wilcke JT, Kok-Jensen A, Dirksen A. Risk of hospital 
admission for obstructive pulmonary disease in α1-antitrypsin 
heterozygotes of phenotype PIMZ. Am J Respir Crit Care Med 
27 Silverman EK, Province MA, Campbell EJ, Pierce JA, Rao DC. Family study of 
α1-antitrypsin deficiency: effects of cigarette smoking, measured genotype, 
and their interaction on pulmonary function and biochemical traits. Genet 
28 Sandford AJ, Chagani T, Weir TD, Connett JE, Anthonisen NR, Pare PD. 
Susceptibility genes for rapid decline of lung function in the Lung Health Study. 
Am J Respir Crit Care Med 2001;163:469–73.
29 Curiel DT, Holmes MD, Okuyama H, Brantly ML, Vogelmeier C, Travis WD, 
Stier LE, Parks WH, Crystal RG. Molecular basis of the liver and lung disease 
associated with the alpha-1-antitrypsin deficiency allele Mvarallo. J Biol Chem 
30 Eriksson S. Alpha-1 antitrypsin deficiency: natural course and therapeutic 
31 Hutchison DCS. α1-Antitrypsin deficiency in Europe: geographical distribution 
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