Deleterious ABCA7 mutations and transcript rescue mechanisms in early-onset Alzheimer’s disease

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Neuropathological description S1: neuropathological examination in an ABCA7 PTC carrier

Neuropathological examination was performed in patient EOD-P1, showing high-level AD neuropathological changes (A3B3C3) as well as a cerebral amyloid angiopathy. The severe global cerebral atrophy was most pronounced in the anterior temporal lobes, with marked neuronal loss and gliosis in the neocortex (especially the frontal, temporal and parietal cortices) and limbic system. Amyloid pathology was frequent in the neocortex, hippocampus, amygdala, cingulum, striatum, and to a lesser extent in the thalamus, cerebellum and brainstem colliculi (CERAD neuritic plaque score of C and a Thal phase of 5, Figure 2a), in addition to abundant amyloid depositions in the wall of the cerebral and cerebellar leptomeningeal vessels (Figure 2b). Tau-positive neurofibrillary tangles and neuropil threads were frequent in the neocortex of the frontal, temporal, parietal and occipital lobes, as well as the cingulum, hippocampus and parahippocampal region (Figure 2c), amygdala, nucleus basalis of Meynert, and to a lesser extent in the brainstem neurons, especially those in the substantia nigra, periaqueductal gray matter, locus caeruleus and raphe nuclei. No α-synuclein immunoreactivity was found.
Method S1: Western blotting

Fresh frozen brain tissue was available for carriers of ABCA7 PTC mutations c.67-1G>A and p.Leu1403fs. Hippocampus was extracted from these two carriers as well as from fresh frozen brain tissue from three AD patients not carrying an ABCA7 PTC mutation. To quantify ABCA7 expression with Western blotting, we adapted the protocol described by Allen and colleagues [1]. Protein lysates were prepared for Western blot using a 0.1% triton lysis buffer (150 mM NaCl, 50 mM Tris pH 7.5, 0.1% Triton). The protein concentration was determined with a BCA assay (Pierce, Rockford, IL, USA), and equal amounts of protein (80µg) were separated on an NuPAGE 3%–8% Tris-Acetate gel and transferred to a PVDF membrane (Hybond P, Amersham Biosciences, Little Chalfont, UK). Membranes were blocked in 5% milk in Phosphate-buffered saline with 0.1% Tween 20 (PBST) and probed overnight at 4°C with the ABCA7 primary antibody designed to epitope aa 2096-2146 (LS-C291064, LifeSpan BioSciences, Seattle, WA, USA; 1/500). Blots were incubated with rabbit IgG horseradish peroxidase–linked secondary antibody for 1 hour. Immunodetection was performed with the ECL-plus chemiluminescent detection system (Amersham Biosciences). Equivalent sample loading was confirmed by probing with anti-glyceraldehyde 3-phosphate dehydrogenase (GAPDH) antibody (GTX100118, Genetex, Irvine, CA, USA; 1/20.000). Protein bands were quantified using ImageQuantTL software, and relative amounts of ABCA7 protein were determined. We observed lower protein expression in PTC mutation carriers in comparison to non-carrier patients (figure S11), confirming a reduction of expression by PTC mutations. Both PTC mutation carriers had no (c.67-1G>A), or low (4%; p.Leu1403fs) potential transcript rescue mechanisms. While a clear difference is shown between carriers and non-carriers, variability in expression is also observed reflective of external modifiers of ABCA7 expression. Of note, this antibody targets the C-terminus of ABCA7, and therefore measures full-length protein. Truncated proteins may be formed, but cannot be observed.
## Supplemental tables

### Table S1: Cohort characteristics for each country of origin

| Country of origin | Patients (n=928) | Controls (n=980) |
|-------------------|------------------|-----------------|
| **Spain**         |                  |                 |
| n=403             |                  | n=223           |
| 58.7% female      |                  | 74.4% female    |
| AAO = 57.8 ± 4.9 years |              | AAI = 58.6 ± 12.8 years |
| APOE ε4-positive = 52.5% |          | APOE ε4-positive = 16.4% |
| **Italy**         |                  |                 |
| n=159             |                  | n=304           |
| 66.0% female      |                  | 59.5% female    |
| AAO = 55.9 ± 7.1 years |              | AAI = 65.3 ± 9.3 years |
| APOE ε4-positive = 38.4% |          | APOE ε4-positive = 23.0% |
| **Sweden**        |                  |                 |
| n=160             |                  | n=295           |
| 63.1% female      |                  | 61.0% female    |
| AAO = 57.9 ± 4.6 years |              | AAI = 64.1 ± 5.4 years |
| APOE ε4-positive = 68.8% |          | APOE ε4-positive = 31.9% |
| **Germany**       |                  |                 |
| n=83              |                  | n=0             |
| 51.8% female      |                  |                 |
| AAO = 58.4 ± 4.7 years |              |                 |
| APOE ε4-positive = 50.6% |          |                 |
| **Portugal**      |                  |                 |
| n=66              |                  | n=120           |
| 59.1% female      |                  | 68.4% female    |
| AAO = 56.3 ± 7.0 years |              | AAI = 66.3 ± 6.1 |
| APOE ε4-positive = 42.4% |          | APOE ε4-positive = 24.2% |
| **Greece**        |                  |                 |
| n=52              |                  | n=35            |
| 61.5% female      |                  | 77.1% female    |
| AAO = 57.5 ± 4.5 years |              | AAI = NA        |
| APOE ε4-positive = 46.2% |          | APOE ε4-positive = 31.4% |
| **Czech Republic** |                  |                 |
| n=5               |                  | n=3             |
| 40.0% female      |                  | 33.3% female    |
| AAO = 52.3 ± 10.5 years |              | AAI = 56.3 ± 10.1 years |
| APOE ε4-positive = 60.0% |          | APOE ε4-positive = 33.3% |

AAI: age at inclusion, AAO: age at onset, NA: not available
Table S2: MinION ABCA7 cDNA sequencing experimental setup.

| Mutation        | Primers                                                      | RNA source | Read depth | cDNA amplicon size (bp) |
|-----------------|--------------------------------------------------------------|------------|------------|-------------------------|
| c.67-1G>A       | CGTTGTCCCTGACCTCTCTCTGTC                                       | brain      | 1488       | 300                     |
|                 | GTCAGCTGCGAAACAG                                             |            |            |                         |
| p.Met370fs      | AACCGGACCTTCCGAGGAG                                          | blood      | 2311       | 398                     |
|                 | TCAGGCTCCAAAGAAGACGAC                                        |            |            |                         |
| p.Glu709fs      | GCCTGGATCTACTCCGTGAC                                         | lymphoblast| 4737       | 649                     |
|                 | AGCTCCTCCGAAAAAGGAAAA                                       |            |            |                         |
| c.3577+1G>C     | CTGCGGACACAGATATGGAG                                          | blood      | 1830       | 296                     |
|                 | AAAGAGGGCAGAGCACAC                                            |            |            |                         |
| p.Trp1336*      | TTTCTGTCTGTGCTGATATTGCCATGTACGGTGCTCAGGTGT                   | lymphoblast| 3279       | 559                     |
|                 | ACTTGCTGTCTGCTCTATCTCCGAGGTAGGTTTTCAGGAG                     |            |            |                         |
| p.Leu1403fs     | TTTCTGTCTGTGCTGATATTGCCATGTACGGTGCTCAGGTGT                   | brain      | 3477       | 559                     |
|                 | ACTTGCTGTCTGCTCTATCTCCGAGGTAGGTTTTCAGGAG                     |            |            |                         |
| c.5570+5G>C     | TGGTGTGGTGCTGGAGAAGACTTG                                      | lymphoblast| 3854       | 420                     |
|                 | GTTTGCTCCCTCCGCTGAG                                           |            |            |                         |

PTC mutations studied on transcript level are denoted by their HGVS notation. Forward and reverse primers were used to amplify the cDNA region of interest. Mutations p.Trp1336* and p.Leu1403fs are in close proximity and the same, but barcoded PCR amplicon, was used. RNA source corresponds to the mutation carrying patient biomaterial. Read depth is the number of reads obtained at the mutation of interest. The amplicon size is denoted according to the canonical ABCA7 transcript (NM_019112) and may change due to alternative splicing.
Table S3: Characteristics of carriers of ABCA7 PTC mutations

| ABCA7 PTC mutation | ID     | Origin | Gender | Diagnosis                  | Presenting features | APOE | Age  | AAD  | DD  | FH  | Other mutation |
|-------------------|--------|--------|--------|----------------------------|---------------------|------|------|------|-----|-----|----------------|
| c.67-1G>A         | EOD-P1 | Spanish| f      | definite AD                | -                   | 34   | ≤59  | 68   | 9   | F   | -              |
|                   | EOD-P2 | Spanish| f      | probable AD                | amnestic            | 33   | 58   | -    | -   | F   | -              |
|                   | EOD-P3 | Spanish| f      | probable AD                | amnestic            | 33   | 64   | -    | -   | F   | -              |
| p.His44fs         | EOD-P4 | Swedish| f      | probable AD                | amnestic            | 23   | 54   | 66   | 12  | S   | -              |
| c.579+1G>T        | EOD-P5 | Portuguese| m   | probable AD                | amnestic            | 44   | 52   | -    | -   | S   | -              |
| c.302+1G>C        | EOD-P6 | Spanish| m      | probable AD                | amnestic + behavioral| 34   | 42   | 49   | 7   | F   | PSEN1 - p.H163R |
| p.Met370fs        | EOD-P7 | Spanish| f      | probable AD                | amnestic            | 43   | 52   | -    | -   | S   | -              |
| p.Cys659fs        | EOD-P8 | Italian| f      | probable AD                | -                   | 33   | 49   | -    | -   | S   | -              |
| p.Glu709fs        | EOD-P9 | Italian| m      | probable AD                | amnestic            | 33   | 58   | -    | -   | S   | -              |
|                   | EOD-P10 | Spanish| f      | probable AD                | amnestic + language | 33   | 55   | 62   | 7   | S   | -              |
|                   | EOD-P11 | Swedish| f      | probable AD                | amnestic            | 34   | 59   | -    | -   | -   | -              |
|                   | EOD-P12 | Spanish| f      | probable AD                | amnestic            | 43   | 60   | -    | -   | F   | -              |
|                   | EOD-P13 | Spanish| f      | probable AD                | amnestic            | 33   | 62   | -    | -   | S   | -              |
|                   | EOD-P14 | Spanish| f      | probable AD                | amnestic            | 33   | 60   | 70   | 10  | F   | -              |
| p.Glu712*         | EOD-P15 | Greek  | f      | AD                          | -                   | 33   | 55   | -    | -   | -   | -              |
| p.Gln732*         | EOD-P16 | Portuguese| m   | probable AD                | amnestic + dysexecutive| 33   | 50   | 57   | 7   | F   | -              |
| p.Thr849fs        | EOD-P17 | Portuguese| f   | probable AD                | amnestic            | 33   | 59   | -    | -   | F   | -              |
|                   | EOD-P18 | Portuguese| f   | probable AD                | amnestic            | 33   | 65   | -    | -   | F   | -              |
| c.3577+1G>C       | EOD-P19 | Spanish| f      | probable AD                | logopenic PPA       | 44   | 61   | 68   | 7   | F   | SORL1 - p.H1813R |
| p.Leu1403fs       | EOD-P20 | Spanish| f      | probable AD                | amnestic            | 34   | 48   | 58   | 10  | S   | -              |
|                   | EOD-P21 | Italian| f      | probable AD                | -                   | 34   | 60   | -    | -   | F   | -              |
|                   | EOD-P22 | Spanish| f      | probable AD                | amnestic            | 44   | 65   | -    | -   | F   | -              |
|                   | EOD-P23 | German  | f      | probable AD                | -                   | 44   | 63   | -    | -   | F   | -              |
|                   | EOD-P24 | Spanish| f      | AD                          | amnestic + slow evolution| 34   | 60   | -    | -   | F   | -              |
| p.Trp1461*        | EOD-P25 | Italian| m      | probable AD                | amnestic + language + dysexecutive| 44   | 57   | -    | -   | F   | -              |
| p.Arg1489*        | EOD-P26 | Spanish| m      | probable AD                | amnestic            | 33   | 55   | -    | -   | F   | -              |
|                   | EOD-P27 | Portuguese| f   | probable AD                | amnestic            | 33   | 55   | -    | -   | S   | -              |
|                   | EOD-P28 | Spanish| m      | AD vs FTD                  | amnestic            | 33   | 50   | -    | -   | S   | -              |
| p.Glu709fs        | EOD-C1 | Italian | m      | CON                         | -                   | 23   | 67   | -    | -   | -   | -              |
| p.Trp1336*        | EOD-C2 | Swedish | f      | CON                         | -                   | 23   | 60   | -    | -   | -   | -              |
|                   | EOD-C3 | Italian | m      | CON                         | -                   | 34   | 58   | -    | -   | -   | -              |
PTC mutations validated in early onset patients (AD) and control individuals (CON). AAD = age at death, AD = Alzheimer’s disease, CON = control, DD = disease duration, FH = familial history, F = familial, PPA = primary progressive aphasia S = sporadic. Age refers to the onset age for patients and inclusion age for controls. Mutation nomenclature is provided according to Human Genome Variation Society (HGVS) on either the transcript or protein (if applicable) level. † This individual carried 2 PTC mutations that segregated on the same haplotype (see Figure S1). § For a description, see Neuropathological description S1. φ This SORL1 missense mutation of unknown importance was previously observed [3]. φ Note: EOD-P6.1 was not included in genetic association testing due to a pathogenic PSEN1 mutation.

| Mutation  | EOD | Country | Gender | Age | Sex | Sであること | AAD | AD | FH | DD | PPA | S | S disease |
|-----------|-----|---------|--------|-----|-----|-------------|-----|----|----|----|-----|---|-----------|
| p.Leu1403fs | EOD-C4 | Italian | f | CON | - | 33 | 44 | - | - | - | - |
| c.4416+2T>G | EOD-C5 | Italian | m | CON | - | 33 | 72 | - | - | - | - |
|           | EOD-C6 | Italian | f | CON | - | 33 | 73 | - | - | - | - |
Table S4: Haplotype of shared ABCA7 PTC mutation carriers

| Marker | Genomic location | Distance to ABCA7 (kb) | Heterozygosity/MAF | p.Glu709fs | p.Leu1409fs |
|--------|------------------|------------------------|--------------------|-------------|-------------|
| D19S814 | chr19:599821-600026 | -440 | NA | - | - |
| D19S886 | chr19:998643-998960 | -41 | 0.63 | 210* | 212* |
| rs3764645 | chr19:1042809 | 0 | 0.4 | G | G |
| rs3764648 | chr19:1044753 | 0 | 0.283 | C | C |
| rs3752234 | chr19:1047002 | 0 | 0.421 | A | G |
| rs3752237 | chr19:1047161 | 0 | 0.305 | G | G |
| rs4147913 | chr19:1049165 | 0 | 0.407 | C | C |
| rs3752240 | chr19:1051214 | 0 | 0.289 | A | G |
| rs3764651 | chr19:1051751 | 0 | 0.49 | G | A |
| rs3764652 | chr19:1052005 | 0 | 0.379 | T | C |
| rs3829687 | chr19:1053299 | 0 | 0.477 | T | C |
| rs3752242 | chr19:1053677 | 0 | 0.382 | A | G |
| rs3752243 | chr19:1054060 | 0 | 0.464 | G | A |
| rs3745842 | chr19:1055191 | 0 | 0.391 | A | G |
| rs3829687 | chr19:1053299 | 0 | 0.477 | T | C |
| rs3752242 | chr19:1053677 | 0 | 0.382 | A | G |
| rs3752243 | chr19:1054060 | 0 | 0.464 | G | A |
| rs3745842 | chr19:1055191 | 0 | 0.391 | A | G |
| rs3829687 | chr19:1053299 | 0 | 0.477 | T | C |
| rs3752242 | chr19:1053677 | 0 | 0.382 | A | G |
| rs3752243 | chr19:1054060 | 0 | 0.464 | G | A |
| rs3745842 | chr19:1055191 | 0 | 0.391 | A | G |
| rs3829687 | chr19:1053299 | 0 | 0.477 | T | C |
| rs3752242 | chr19:1053677 | 0 | 0.382 | A | G |
| rs3752243 | chr19:1054060 | 0 | 0.464 | G | A |
| rs3745842 | chr19:1055191 | 0 | 0.391 | A | G |
| rs3829687 | chr19:1053299 | 0 | 0.477 | T | C |
| rs3752242 | chr19:1053677 | 0 | 0.382 | A | G |
| rs3752243 | chr19:1054060 | 0 | 0.464 | G | A |
| rs3745842 | chr19:1055191 | 0 | 0.391 | A | G |
| rs2242437 | chr19:1065563 | 0 | 0.395 | G | C |
| D19S814 | chr19:1413756-1414123 | 348 | 0.73 | - | - |
| D19S886 | chr19:2359696-2359954 | 1294 | 0.82 | - | - |
| D19S424 | chr19:3226372-3226704 | 2161 | 0.79 | - | - |
| D19S814 | chr19:4392405-4392667 | 3327 | 0.78 | - | - |
| D19S177 | chr19:5517296-5517670 | 4452 | 0.76 | - | - |

For individuals sharing the same PTC mutations we determined haplotypes based on SNPs and STR markers spanning ABCA7 and flanking regions as previously described [2]. Briefly, we selected 18 common SNPs (rs-numbers) that were covered within our targeting assay while passing Hardy Weinberg Equilibrium quality control (p>0.001). In addition, seven STR markers (D19S-notation) were genotyped with FAM-labeled multiplex PCR, supplemented with GeneScan™ 500 LIZ™ size standard (Thermo Fisher Scientific, Waltham, USA) after which capillary fragment analysis was performed on an ABI 3730 DNA Analyzer (Thermo Fisher Scientific, Waltham, USA). Heterozygosities of STR markers were obtained from
the Marshfield Clinic database (www.marshfieldclinic.org) and MAF of SNPs was based on 1000 genomes data. *ABCA7* is located in the 19p subtelomeric region, hence only two STR markers were selected on the telomeric side of *ABCA7*. The shared haplotype is shown per PTC variant. ‘-’ = Samples did not share this marker. ‘*’This STR allele was absent in 1 individual carrying the respective mutation.
Table S5: Carriers of *ABCA7* c.5570+5G>C in the EU EOD consortium

| Sample ID | Status | Gender | Origin  | APOE | Age | Family History | c.5570+5G>C zygosity |
|-----------|--------|--------|---------|------|-----|----------------|----------------------|
| EOD-P45   | AD     | m      | Italian | 33   | 51  | S              | Heterozygous         |
| EOD-P46   | AD     | m      | Spanish | 34   | 53  | S              | Heterozygous         |
| EOD-P47   | AD     | m      | Spanish | 44   | 61  | F              | Heterozygous         |
| EOD-P48   | AD     | m      | Swedish | 33   | 56  |                | Homozygous           |
| EOD-P49   | AD     | f      | Swedish | 33   | 56  |                | Heterozygous         |
| EOD-P50   | AD     | f      | Swedish | 33   | 64  |                | Heterozygous         |
| EOD-C16   | CON    | f      | Italian | 33   | -   |                | Heterozygous         |
| EOD-C17   | CON    | -      | Portuguese | 33  | -   |                | Heterozygous         |
| EOD-C18   | CON    | m      | Spanish | 34   | -   |                | Heterozygous         |
| EOD-C19   | CON    | f      | Swedish | 33   | 61  |                | Heterozygous         |
| EOD-C20   | CON    | f      | Swedish | 23   | 61  |                | Heterozygous         |
| EOD-C21   | CON    | f      | Swedish | 33   | 66  |                | Heterozygous         |
| EOD-C15   | CON    | f      | Swedish | 33   | 60  |                | Heterozygous         |

Age = onset age for patients (AD) and inclusion age for controls (CON). F = positive familial history (affected first degree relative). S = sporadic. Heterozygous = GC genotype, Homozygous = CC genotype.
Table S6: Carriers of ABCA7 missense mutations with predicted deleterious effects (CADD > 20)

| Mutation coordinates and nomenclature | Effect prediction | Carrier characteristics |
|---------------------------------------|-------------------|-------------------------|
| Genomic position | HGVS (coding)  | HGVS (protein) | PolyPhen category | SIFT category | Phred CADD score | Sample ID | Status | Gender | Origin | APOE | Age | FH |
| chr19:1045025 | c.1240G>A | p.Ala414Thr | benign | deleterious | 24.1 | EOD-P30 | AD | m | German | 33 | 59 | - |
| chr19:1046288 | c.1505G>C | p.Gly502Ala | possibly damaging | deleterious | 25 | EOD-C14 | CON | m | Swedish | 33 | 60 | - |
| chr19:1047168 | c.1858C>T | p.Leu620Phe | probably damaging | deleterious | 31 | EOD-P32 | AD | f | Greek | 34 | 60 | - |
| chr19:1047168 | c.1858C>T | p.Leu620Phe | probably damaging | deleterious | 31 | EOD-P36 | AD | f | Italian | 34 | 58 | F |
| chr19:1047169 | c.1859T>C | p.Leu620Pro | probably damaging | deleterious | 28.6 | EOD-P41 | AD | f | Spanish | 44 | 57 | F |
| chr19:1047336 | c.2026G>A | p.Ala676Thr | probably damaging | deleterious | 27.1 | EOD-P40† | AD | f | Spanish | 33 | 58 | F |
| chr19:1047336 | c.2026G>A | p.Ala676Thr | probably damaging | deleterious | 27.1 | EOD-C10† | CON | m | Portuguese | 33 | - | - |
| chr19:1047336 | c.2026G>A | p.Ala676Thr | probably damaging | deleterious | 27.1 | EOD-C11† | CON | f | Spanish | 33 | - | - |
| chr19:1047498 | c.2114C>A | p.Ala705Asp | probably damaging | deleterious | 28.9 | EOD-P37 | AD | f | Spanish | 33 | 56 | S |
| chr19:1049283 | c.2399C>T | p.Pro800Leu | possibly damaging | tolerated | 24.7 | EOD-P31 | AD | f | Greek | 33 | 49 | - |
| chr19:1051006 | c.2639G>T | p.Arg880Gln | probably damaging | deleterious | 34 | EOD-P29 | AD | f | German | 44 | 58 | - |
| chr19:1051006 | c.2639G>T | p.Arg880Gln | probably damaging | deleterious | 34 | EOD-P44 | AD | m | Swedish | 34 | 59 | - |
| chr19:1051006 | c.2639G>T | p.Arg880Gln | probably damaging | deleterious | 34 | EOD-C13 | CON | f | Swedish | 34 | 61 | - |
| chr19:1051944 | c.2966G>T | p.Arg989His | probably damaging | deleterious | 33 | EOD-C15 | CON | f | Spanish | 33 | 59 | - |
| chr19:1051944 | c.2966G>T | p.Arg989His | probably damaging | deleterious | 33 | EOD-C12 | CON | f | Italian | 33 | 60 | - |
| chr19:1051964 | c.2986C>T | p.His996Tyr | probably damaging | deleterious | 28.1 | EOD-C7 | CON | m | Italian | 33 | - | - |
| chr19:1051964 | c.2986C>T | p.His996Tyr | probably damaging | deleterious | 28.1 | EOD-C8 | CON | m | Italian | 33 | - | - |
| chr19:1052067 | c.3089G>T | p.Gly1030Val | possibly damaging | deleterious | 26.9 | EOD-P39 | AD | f | Spanish | 34 | 65 | S |
| Chr     | Mutation Type | Mutation | p.Amino Acid | Functional Implication | Age | FH | Gender | Ethnicity | Onset Age | Inclusion Age | Notes |
|---------|---------------|----------|--------------|-----------------------|-----|----|--------|-----------|-----------|--------------|-------|
| chr19:1053504 | c.3397G>A     | p.Gly1133Arg | probably damaging | deleterious             | 34  | f  | Italian | 34        | 62        | F            |       |
| chr19:1057919 | c.4886C>T     | p.Ser1629Leu | probably damaging | deleterious             | 35  | f  | Spanish | 33        | 50        | S            |       |
| chr19:1057949 | c.4922_4924delTCT | p.Phe1641del | NA           | NA                     | 20.7| EOD-P40† | AD | f  | Spanish | 33        | 58        | F            |       |
| chr19:1058635 | c.5168C>T     | p.Ser1723Leu | probably damaging | deleterious             | 33  | EOD-P40† | AD | f  | Spanish | 33        | 58        | F            |       |
| chr19:1058635 | c.5168C>T     | p.Ser1723Leu | probably damaging | deleterious             | 33  | EOD-P42  | AD | f  | Spanish | 33        | 47        | -            |       |
| chr19:1058635 | c.5168C>T     | p.Ser1723Leu | probably damaging | deleterious             | 33  | EOD-C10† | CON | m  | Portuguese | 33       | -         | -            |       |
| chr19:1058939 | c.5206G>T     | p.Ala1736Ser | possibly damaging | deleterious             | 25.7| EOD-P33  | AD | f  | Italian | 33        | 62        | -            |       |
| chr19:1059079 | c.5458G>A     | p.Gly1820Ser | probably damaging | deleterious             | 28.5| EOD-C9   | CON | m  | Italian | 33        | 46        | -            |       |
| chr19:1061817 | c.5500A>G     | p.Thr1834Ala | possibly damaging | deleterious             | 26  | EOD-P35  | AD | f  | Italian | 33        | 47        | S            |       |

Individuals carrying a missense mutation with a Phred-scaled CADD score higher than 20 are shown. In addition, predicted effects of PolyPhen and Sift (incorporated into CADD) are shown. Age = onset age for patients (AD) and inclusion age for controls (CON). FH = Familial History. F = positive familial history (affected first degree relative). S = sporadic. † These individuals carried a “double deleterious missense” haplotype.
Table S7: Meta-analysis of association between common coding ABCA7 SNPs and EOAD in the EU EOD consortium.

| Genomic position (hg19) | Mutation coordinates and nomenclature | HGVS (coding) | HGVS (protein) | dbSNP | MAF | AD (%) | CON (%) | OR (95% CI) | p-value |
|------------------------|--------------------------------------|---------------|----------------|-------|-----|--------|---------|-------------|--------|
| chr19:1043103          | c.G643A                              | p.G215S       | rs72973581A    | A     | 3.33| 4.74   |         | 0.60 (0.42 - 0.87) | 0.006  |
| chr19:1061804          | c.T5487C                             | p.N1829N      | rs78320196C    | C     | 3.74| 5.68   |         | 0.65 (0.47 - 0.90) | 0.009  |
| chr19:1042809          | c.A563G                              | p.E188G       | rs3764645G     | G     | 41.40| 45.32  |         | 0.86 (0.75 - 0.99) | 0.030  |
| chr19:1064193          | c.G5985A                             | p.L1995L      | rs4147930G     | G     | 27.93| 30.04  |         | 0.89 (0.76 - 1.03) | 0.121  |
| chr19:1053524          | c.C3417G                             | p.E188G       | rs3764645G     | G     | 41.40| 45.32  |         | 0.86 (0.75 - 0.99) | 0.030  |
| chr19:1051214          | c.A2745G                             | p.V915V       | rs3752240G     | G     | 32.78| 35.28  |         | 0.91 (0.79 - 1.06) | 0.218  |
| chr19:1043748          | c.A955G                              | p.T319A       | rs3752232G     | G     | 2.56 | 3.17   |         | 0.77 (0.50 - 1.20) | 0.256  |
| chr19:1047537          | c.A2153C                             | p.N718T       | rs3752239C     | C     | 2.56 | 3.23   |         | 0.84 (0.55 - 1.28) | 0.424  |
| chr19:1041852          | c.G183T                              | p.L61L        | rs3764644T     | T     | 2.76 | 3.31   |         | 0.86 (0.57 - 1.29) | 0.459  |
| chr19:1054060          | c.A3528G                             | p.L1176L      | rs3752243G     | G     | 43.16| 41.79  |         | 1.05 (0.92 - 1.21) | 0.452  |
| chr19:1052005          | c.C3027T                             | p.A1009A      | rs3764652T     | T     | 43.43| 42.18  |         | 1.05 (0.92 - 1.21) | 0.458  |
| chr19:1062164          | c.SS71-7T>C                          | -             | rs4147920C     | C     | 2.60 | 3.18   |         | 0.88 (0.57 - 1.35) | 0.566  |
| chr19:1062192          | c.T5592C                             | p.A1864A      | rs4147921C     | C     | 2.55 | 3.07   |         | 0.88 (0.57 - 1.36) | 0.571  |
| chr19:1056065          | c.A4239G                             | p.R1413R      | rs881768G      | G     | 43.08| 41.98  |         | 1.05 (0.91 - 1.20) | 0.516  |
| chr19:1058176          | c.A5057G                             | p.Q1686R      | rs4147918G     | G     | 2.71 | 3.42   |         | 0.89 (0.59 - 1.34) | 0.582  |
| chr19:1044712          | c.A1184G                             | p.H395R       | rs3764647G     | G     | 2.98 | 3.72   |         | 0.89 (0.58 - 1.36) | 0.586  |
| chr19:1055191          | c.G4046A                             | p.R1349Q      | rs3745842A     | A     | 42.52| 41.45  |         | 1.03 (0.90 - 1.19) | 0.652  |
| chr19:1047161          | c.A1851G                             | p.G617G       | rs3752237A     | A     | 41.50| 40.48  |         | 1.02 (0.88 - 1.19) | 0.768  |
| chr19:1056492          | c.G4580C                             | p.G1527A      | rs3752246G     | G     | 21.48| 20.52  |         | 1.02 (0.86 - 1.20) | 0.850  |
| chr19:1049269          | c.G2385A                             | p.L795L       | rs4147914A     | A     | 16.18| 16.60  |         | 0.98 (0.81 - 1.17) | 0.799  |
| chr19:1049305          | c.C2421A                             | p.V807V       | rs4147915A     | A     | 12.72| 12.60  |         | 1.02 (0.83 - 1.26) | 0.823  |
| chr19:1047002          | c.A1824G                             | p.A608A       | rs3752234A     | A     | 46.68| 46.37  |         | 0.99 (0.86 - 1.14) | 0.884  |

Genomic coordinates are based on hg19. Fixed effects (Cochran-Mantel-Haenszel) meta-analysis was performed on all common (minor allele frequency (MAF) > 1%) coding SNPs. HGVS = mutation nomenclature according to the Human Genome Variation Society. dbSNP notations refer to Reference SNP IDs (rs) from dbSNP build 142. MA = minor allele. AD = Alzheimer’s Disease. CON = Control individual. Odds ratios (OR) and 95% confidence intervals (CI) are calculated for the MA. The study-wide multiple testing corrected p-value cutoff is 0.0033.
Supplemental figures

**Figure S1: Genotype phasing of two PTC mutations in one individual.** One patient (EOD-P5) carried two PTC mutations (c.206G>A (p.Trp69*) and c.579+1G>T). Allele-specific PCR was performed to resolve haplotype phase. The final 3’ nucleotide of the reverse primer corresponded with either the G (5’-GGATGAGTGGGGCTCGTAC-3’) or T (5’-GGATGAGTGGGGCTCGTAA-3’) allele of variant c.579+1G>T. PCR amplification of the fragment containing c.206G>A was separately performed for each c.579+1G>T allele in combination with a common forward primer (5’-CAGGGACCAGGCACTTTGTG-3’), after which Sanger sequencing was performed with the forward primer. Both mutations segregated in cis as shown in the figure: The reference c.579+1G>T[G]-allele (upper chromatogram panel) carried the reference c.206G>A[G]-allele, while the PTC c.579+1G>T[T]-allele (second chromatogram panel) co-segregated with the PTC c.206G>A[A]-allele. The bottom nucleotide track corresponds to the reference sequence (hg19).
Figure S2: Dot plot showing the onset ages of ABCA7 PTC carriers versus carriers of a pathogenic PSEN1, PSEN2 or APP mutation. Mann-Whitney U test p value 0.0002. The cross denotes an individual who carries an ABCA7 PTC mutation (c.302+1G>C) in addition to a pathogenic PSEN1 p.His163Arg mutation. Note: all are EOAD patients, hence all onset ages are equal to or below 65 years.
**Figure S3: MinION and Sanger cDNA sequencing of ABCA7 p.Met370fs.** The cDNA sequencing shown here, was derived from blood RNA of a p.Met370fs (c.1109dupT) mutation carrier. The upper panels (a-d) correspond to IGV visualization of MinION sequencing data: (a) Coordinates on chromosome 19 (hg19) and location of the PTC mutation (red arrow), corresponding to features in panels b, c, and d. (b) Coverage plot in which the height of each bar corresponds to the read depth at that nucleotide position. Gray color indicates that the hg19 reference nucleotide was observed (c) A snapshot of sequencing reads. Each gray bar represents aligned sequences, which are connected by blue lines that correspond to splicing events. In this example, three reads show skipping of exon 11 (which contains p.Met370fs). (d) A color code corresponding to reference nucleotides is shown and below blue bars and lines depict ABCA7 exons and introns, respectively. The red lines show the region that is represented in more detail below (e-f) using Sanger sequencing: (e) The reference sequence. (f) Forward (top) and reverse (bottom) Sanger sequencing of the c.1109dupT mutation (red arrow), which causes insertion of a T allele and hence a shift in nucleotides (yellow highlights).
RNA and cDNA were obtained from patient lymphoblasts. Sequencing reads were visualized with IGV. 

(a) Coordinates on chromosome 19 (hg19) and location of the PTC mutation (red arrow), corresponding to features in panels b, c, d, and e. (b) Coverage plot in which the height of each bar corresponds to the read depth at that nucleotide position. Colors are shown depending on the observed nucleotides, and gray color indicates that the hg19 reference nucleotide was observed. (c) This panel represents MinION cDNA sequencing. Gray bars correspond to aligned sequences and blue lines denote splicing events. Hence several canonically spliced transcripts are shown, as well as reads skipping exon 30 (4 reads in this figure). Due to long length sequencing on a MinION platform, each read shown here encompasses the entire selected part of the ABCA7 transcript. The MinION produces a higher error rate than Illumina as depicted by the mismatches, deletions, and insertions shown in the aligned sequences. Accuracy, however, is high enough for alignment and calculation of splicing, and at high read depth, these errors are negated. (d) This panel shows RNAseq, validating the exon skipping shown in panel c. Read lengths (depicted by gray segments) produced by Illumina RNAseq are much shorter than those generated by MinION. Furthermore the number of reads in RNAseq aligning to ABCA7 is very low due to low
expression of this gene. Even though the error rate is lower this compromises a semi-quantitative assessment of alternative splicing isoforms. (e) An overview of the nucleotides (color code track) and ABCA7 gene layout (blue). The red lines denote a region that was analyzed at higher depth in panels f and g. (f) Zoomed in RNAseq reads clearly depict the presence of the nonsense p.Trp1336[A]-allele (red arrow), as well as exon 30 skipping. (g) The reference nucleotides and amino acids to which the reads in panel f align.
Figure S5: MinION sequencing of ABCA7 p.Leu1403fs brain cDNA tissue. IGV visualization of long-read cDNA sequencing. (a) Coordinates on chromosome 19 (hg19) and location of the PTC mutation (red arrow), corresponding to features in panels b, c, and d. (b) Coverage plot in which the height of each bar corresponds to the read depth at that nucleotide position. Colors are shown depending on the observed nucleotides, and gray color indicates that the hg19 reference nucleotide was observed. (c) Each row represents aligned sequencing reads (gray bars) and splicing events (blue lines). Skipping of exon 31 harboring the p.Leu1403fs mutation is visible (4%). This visualization also illustrates relatively common skipping of exon 30 (30%). Of note, one of the other PTC mutations, p.Trp1336* (observed in a separate control individual – see Figure S4), is located in this exon. (d) An overview of the nucleotides (color code track) and ABCA7 gene layout. The red lines denote PTC carrying exon 31, which is shown in more detail in panels e and f. (e) A snapshot of reads covering exon 31, which depicts the deletion (black horizontal lines) of a thymidine residue (red arrow) leading to p.Leu1403fs. (f) Reference nucleotides and amino acids in exon 31.
Figure S6: Validation of ABCA7 exon 11 skipping transcripts with Illumina RNAseq. IGV visualization of sequencing reads originating from lymphoblast RNA [3]. Coordinates on chromosome 19 (hg19) are shown on top. The three panels each correspond to a different individual and depict aligned sequencing reads (gray bars) as well as splicing events (blue connecting lines). At the bottom, a color code track corresponds to different nucleotides in the reference sequence and the canonical ABCA7 transcript (NM_019112) is shown in blue.

MinION sequencing (Figure S3) demonstrated inframe exon 11 skipping as a potential rescue mechanism for PTC mutations located in exon 11 (e.g. p.Met370fs). The RNAseq samples depicted here, do not carry a PTC mutation in exon 11, and show a common occurrence of this novel ABCA7 isoform (red arrows). Exon 11 skipping is therefore not exclusive to exon 11 PTC mutation carriers, but can potentially rescue the negative effect of a PTC mutation in exon 11.
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Figure S7: Validation of *ABCA7* exon 30 and 31 skipping with Illumina RNAseq. IGV visualization of sequencing reads originating from lymphoblast RNA [3]. Coordinates on chromosome 19 (hg19) are shown on top. The three panels below each correspond to a different individual and depict aligned sequencing reads (gray bars) as well as splicing events (blue connecting lines). At the bottom, a color code track corresponds to different nucleotides in the reference sequence and the canonical *ABCA7* transcript (NM_019112) is shown in blue.

Both exon 30 and 31 contain known PTC mutations (respectively p.Trp1336* and p.Leu1403fs). With MinION cDNA sequencing we observed inframe skipping of exon 30 and 31, which has the potential to diminish deleterious mutational effects. With RNAseq data we can validate both exon skipping events (red arrows). The upper panel corresponds to an p.Trp1336* carrier (location of the mutation is depicted with a blue arrow).
**Figure S8: MinION sequencing of ABCA7 p.Glu709fs.** IGV visualization of sequencing reads from MinION cDNA sequencing on lymphoblast RNA of a p.Glu709fs carrying patient. (a) Coordinates on chromosome 19 (hg19), corresponding to features in panels b, c, and d. (b) Coverage plot in which the height of each bar corresponds to the read depth at that nucleotide position. Colors are shown depending on the observed nucleotides, and gray color indicates that the hg19 reference nucleotide was observed. (c) aligned sequencing reads (gray bars) and splicing events (blue lines). (d) An overview of the nucleotides (color code track) and ABCA7 gene layout (blue) along with annotated events of interest (red).

The 7bp deletion (black horizontal lines in panel c, and drop in coverage in panel b) leading to p.Glu709fs is observed in ABCA7 transcripts. While most reads follow a canonical splicing pattern, some p.Glu709fs transcripts are alternatively spliced by usage of a cryptic splice donor site 23bp upstream of the canonical splice donor site, with restoration of the reading frame as a consequence. Additionally other splicing events are observed as well (e.g. usage of a cryptic exon 16 splice donor site 73bp upstream).
Figure S9: Validation of alternative splicing of ABCA7 exon 16 with Illumina RNAseq. IGV visualization of sequencing reads originating from lymphoblast RNA [3]. Coordinates on chromosome 19 (hg19) are shown on top. The three panels below each correspond to a different individual and depict aligned sequencing reads (gray bars) as well as splicing events (blue connecting lines). At the bottom, a color code track corresponds to different nucleotides in the reference sequence and the canonical ABCA7 transcript (NM_019112) is shown in blue.

Using MinION cDNA sequencing we observed novel ABCA7 isoforms which comprised alternatively spliced exon 16 (Figure S8). One particular splicing event encompassed the usage of a cryptic splice donor site 23bp upstream of the canonical splice donor site. In conjunction with a p.Glu709fs mutation (7bp deletion) this alternative splicing event can restore the transcript reading frame. Here, RNAseq analysis validates the existence of this isoform (red arrows). The upper RNAseq panel, corresponds to a p.Glu709fs carrier and depicts the limitations of RNAseq. No 7bp deletion is observed in exon 16, however, this is most likely a result of the low ABCA7 coverage in RNAseq. Furthermore it cannot be determined whether the alternative splicing observed in this panel is in phase with the mutation.
RNA and cDNA were obtained from c.5570+5G>C carrying patient lymphoblasts. Sequencing results are visualized with IGV. (a) Coordinates on chromosome 19 (hg19) and location of the PTC mutation (red arrow), corresponding to features in panels b, c, d, and e. (b) Coverage plot in which the height of each bar corresponds to the read depth at the corresponding position. Colors are shown depending on the observed nucleotides, and gray color indicates that the hg19 reference nucleotide was observed. (c) This panel represents MinION cDNA sequencing. Gray bars correspond to aligned sequences and blue lines denote splicing events. Several differentially spliced transcripts are observed: exon 41 skipping, 14bp intron 41 retention due to c.5570+5G>C, complete intron 41 retention and intron 41 retention starting from a cryptic splice acceptor site 204bp upstream of the canonical exon 42 splice acceptor. The accuracy of reads generated by the MinION platform, may be lower than Illumina sequencing (e.g. panel d), but is sufficient to align individual reads, which reveal a complex splicing pattern of this region. (d) (Partial) intron 41 retention is validated by RNAseq. This panel shows that to fully comprehend the splicing complexity, long reads (panel b) - instead of the short Illumina reads (gray segments) depicted

**Figure S10: MinION cDNA sequencing and Illumina RNA sequencing of ABCA7 c.5570+5G>C.**

here - provide more clarity. Intron 41 retentions for instance are not spanned by the short reads, and it is therefore unknown whether these reads originate from a full or partial intron retention transcript. (e) An overview of the nucleotides (color code track) and ABCA7 gene layout (blue). The red lines denote a region that was analyzed at higher depth in panels e and f. (f) Zoomed in RNAseq reads confirm the c.5570+5G>C[C]-allele as the cause of 14bp intron retention. (g) The reference nucleotides, amino acids to which the reads in panel e align, and location of c.5570+5G>C (red arrow).
Figure S11: Hippocampal ABCA7 protein quantification in AD patients with or without an ABCA7 PTC mutation. (a) Western blotting (Method S1) was performed on hippocampal brain tissue from three AD patients without an ABCA7 PTC mutation (lanes 1, 2, and 3) and two carriers: c.67-1G>A (lane 4) and p.Leu1403fs (lane 5). A protein band is detected at approximately 234kDa, which corresponds to full length ABCA7 protein (ENSP00000414062). In addition, immunodetection of glyceraldehyde 3-phosphate dehydrogenase (GAPDH) is shown below. (b) Relative quantity (RQ) of ABCA7 is normalized by GAPDH signal strength. Western blotting was repeated twice.
Supplemental references

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