High-throughput imaging of ATG9A distribution as a diagnostic functional assay for adaptor protein complex 4-associated hereditary spastic paraplegia

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Adaptor protein complex 4-associated hereditary spastic paraplegia is caused by biallelic loss-of-function variants in AP4B1, AP4M1, AP4E1 or AP4S1, which constitute the four subunits of this obligate complex. While the diagnosis of adaptor protein complex 4-associated hereditary spastic paraplegia relies on molecular testing, the interpretation of novel missense variants remains challenging. Here, we address this diagnostic gap by using patient-derived fibroblasts to establish a functional assay that measures the subcellular localization of ATG9A, a transmembrane protein that is sorted by adaptor protein complex 4. Using automated high-throughput microscopy, we determine the ratio of the ATG9A fluorescence in the trans-Golgi-network versus cytoplasm and ascertain that this metric meets standards for screening assays (Z-factor robust >0.3, strictly standardized mean difference >3). The ‘ATG9A ratio’ is increased in fibroblasts of 18 well-characterized adaptor protein complex 4-associated hereditary spastic paraplegia patients [mean: 1.54 ± 0.13 versus 1.21 ± 0.05 (standard deviation) in controls] and receiver-operating characteristic analysis demonstrates robust diagnostic power (area under the curve: 0.85, 95% confidence interval: 0.849–0.852). Using fibroblasts from two individuals with atypical clinical features and novel biallelic missense variants of unknown significance in AP4B1, we show that our assay can reliably detect adaptor protein complex 4 function. Our findings establish the ‘ATG9A ratio’ as a diagnostic marker of adaptor protein complex 4-associated hereditary spastic paraplegia.
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**Introduction**

The hereditary spastic paraplegias (HSPs) are a heterogeneous group of over 80 different neurogenetic disorders that share the clinical feature of spasticity due to corticospinal tract dysfunction. Childhood-onset forms of HSP often pose a diagnostic challenge given their non-specific clinical presentation and slow progression in many cases. The increasing availability of next-generation sequencing is beginning to close this diagnostic gap, yet, at the same time, has broadened the phenotypic spectrum associated with HSP genes. As for other rare disorders, the interpretation of genetic variants remains a significant challenge, particularly for missense variants associated with novel or atypical phenotypes. In this setting, functional assays in patient-derived cells can aid the interpretation of variants and help establish a clinical diagnosis. This is supported by the current American College of Medical Genetics and Genomics (ACMG) recommendations for variant classification (criterion PS3), which integrate results from ‘well-established’ functional assays.

Adaptor protein complex 4 (AP-4)-associated hereditary spastic paraplegia (AP-4-HSP) encompasses four forms of childhood-onset HSP, caused by biallelic variants in the subunits of AP-4. This includes AP4B1-associated SPG47 (OMIM #614066), AP4M1-associated SPG50 (OMIM #612936), AP4E1-associated SPG51 (OMIM #613744) and AP4S1-associated SPG52 (OMIM #614067). AP-4 is a member of the adaptor protein complex family of coated vesicle adaptors, which functions in the generation of intracellular transport vesicles carrying transmembrane cargo proteins (recently reviewed in Sanger et al.). Owing to the obligate nature of the heterotetrameric AP-4 complex, loss of AP-4 function is common to all forms of AP-4-HSP. Recent work has shown that AP-4 deficiency leads to prominent changes in the subcellular localization of the AP-4 cargo protein ATG9A, including in patient-derived cells. ATG9A is the only conserved transmembrane autophagy-related protein and in mammalian cells cycles between the trans-Golgi network (TGN) and ATG9A vesicles, which associate with endosomes and autophagosome formation sites. ATG9A is a lipid scramblase important for autophagic vesicle formation and is essential for axonal function, as demonstrated in CNS-specific Atg9a knockout mice.

Informed by this emerging understanding of AP-4 biology, we here set out to validate a high-throughput ATG9A translocation assay in patient-derived fibroblasts as a diagnostic marker of AP-4-HSP. Using an automated microscopy platform that allows us to measure the subcellular localization of ATG9A in thousands of cells in a single run, we show that the ATG9A fluorescence signal in the TGN is robustly increased in well-characterized AP-4-HSP patients. Our findings establish the ‘ATG9A ratio’ as a robust indicator of AP-4 function that can provide a functional validation of novel variants in AP-4 subunit genes.

**Materials and methods**

**Patients and clinical information**

This study was approved by the Institutional Review Board at Boston Children’s Hospital (IRB-P00033016 and IRB-P00016119). Patients with AP-4-HSP were recruited from the AP-4-HSP International Registry (ClinicalTrials.gov Identifier: NCT04712812). All patients underwent clinical exome sequencing showing biallelic variants in AP4B1, AP4M1 or AP4S1. Clinical information was collected systematically using the AP-4-HSP Natural History Study Questionnaire. Severity of motor symptoms and corticospinal tract dysfunction were assessed using the Spastic Paraplegia Rating Scale and a four-stage mobility score. Clinical information of 13 patients has been reported previously.

**Variant interpretation and scoring**

Variants were standardized to the following transcripts: AP4B1 (NM_001253852.3, GRCh37/hg19), AP4M1 (NM_004722.4, GRCh37/hg19), AP4E1 (NM_007347.5, GRCh37/hg19) and AP4S1 (NM_007077.4, GRCh37/hg19). Variants were classified following ACMG guidelines using VarSome and InterVar. For data shown in Figs 1 and 2 and Supplementary Fig. 1, all reported variants from the AP-4-HSP International Registry were...
annotated as described previously,\textsuperscript{22} using the most recent version of all tools and scores.\textsuperscript{23–28}

**Analysis of variant spectrum and missense variant distribution in AP-4 protein domain structures**

All variants from individuals included in the AP-4-HSP International Registry were mapped to the secondary protein structures of the AP-4 subunits derived from UniProt (Supplementary File 1). Next, the available domain information from UniProt\textsuperscript{39} and pfam\textsuperscript{40} were compared with the homology model of the AP-4 complex reported recently.\textsuperscript{31} Domains predicted by pfam showed a greater overlap and were thus chosen for further analyses. For AP4S1, alignment was transferred from isoform 1 and isoform 2 of Q9Y6B7 using InterPro.\textsuperscript{32} Constraint scores of protein regions were analysed using CADD PHRED,\textsuperscript{27} M-CAP and REVEL scores (Supplementary File 1).

![Figure 1 AP4B1 and AP4M1 subunit structures with variant distribution and CADD scores for all possible missense variants.](image-url)

**Figure 1** AP4B1 and AP4M1 subunit structures with variant distribution and CADD scores for all possible missense variants. (A and B) Schematic representation of AP4B1 (based on UniProt Q9Y6B7) and AP4M1 (based on UniProt O00189) proteins with their respective aligned units identified by pfam. Blue dots represent reported missense variants, splice variants are depicted in pink and truncating variants in red. Novel variants are highlighted in grey. Length of vertical lines corresponds to the CADD score for each variant. Also shown are the CADD PHRED (version 1.6) scores across the protein secondary structure for all possible missense variants in AP4B1 and AP4M1 respectively. The red horizontal line marks the recommended cut-off of 20. Similar analyses for M-CAP and REVEL scores are found in Supplementary Fig. 1. Data are summarized in Supplementary File 1.
Reagents and antibodies

The following reagents were used: Bovine serum albumin (AmericanBio, #9048-46-8), saponin (Sigma, #47036-50G-F), Molecular Probes Hoechst 33258 (#H3569) and Alexa Fluor 647-labelled phalloidin (Thermo Fisher Scientific, #A22287). The following primary antibodies were used: Anti-ATG9A at 1:400 (Abcam # ab108338), anti-TGN46 at 1:800 (Bio-Rad #AHP500G), anti-AP4E1 and anti-AP4B1 (both produced in house). Fluorescently labelled secondary antibodies used in this study were all purchased from Thermo Fisher Scientific and used at 1:2000 (#A11008, #A11016, #A21245). Horseradish peroxidase-conjugated secondary antibodies were purchased from Sigma-Aldrich and used at 1:5000 dilution. Lentivirus to express human AP4B1 under a PGK promoter was generated as described previously.11

Fibroblast cell culture

Fibroblast lines were established from routine skin punch biopsies.11 Primary human skin fibroblasts were cultured and maintained as described previously.35 Briefly, cells were maintained in Dulbecco’s Modified Eagle Medium (DMEM) high glucose (Gibco, #11960044) supplemented
with 20% foetal bovine serum (FBS) (Gibco, #10082147), penicillin 100 U/ml and streptomycin 100 μg/ml (Gibco, #15140122). Cells were kept in culture for up to 8 passages and routinely tested for the presence of mycoplasma contamination. For high-throughput imaging shown in Figs 3–5, fibroblasts were seeded onto 96-well plates (Greiner Bio-One, #655090) at a density of $3 \times 10^3$ per well or onto 384-well plates (Greiner Bio-One, #781090) at a density of $2 \times 10^3$ per well using the Multidrop Combi Reagent Dispenser (Thermo Fisher Scientific, #11388-558).

### AP4B1 knockout SH-SY5Y cells

AP4B1 knockout SH-SY5Y cells ($AP4B1^{KO}$) were a mixed population of CRISPR/Cas9-edited cells, as previously described.\(^{10}\) SH-SY5Y cells were maintained in DMEM/F12 (Gibco, Cat# 11320033) supplemented with...
10% heat-inactivated FBS (Gibco, Cat# 10438026), 100 U/ml penicillin and 100 μg/ml streptomycin at 37°C under 5% CO₂. SH-SY5Y cells were differentiated into a neuron-like state by culture in the presence of retinoic acid as described previously. Cells were seeded at 25% confluency in standard culture medium, and the following day
the medium was replaced with neurobasal medium with B27 (Thermo Fisher Scientific, #11320082, #17504-044) containing 10 μM all-trans-retinoic acid (MedChemExpress, #HY-14649). After 48 h, cells were split 1:4 and cultured for 3 further days in the presence of retinoic acid (5 days total).

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**Figure 5** The ATG9A ratio can provide a functional assessment of novel missense variants. (A) Mean ATG9A ratio of all assay plates in AP-4-HSP patients with biallelic loss-of-function variants (P−/−) and two patients with novel biallelic missense variants (PVUS/VUS_01 and PVUS/VUS_02) (one-way ANOVA with Dunnett’s post hoc test, P−/+ versus PVUS/VUS_01, P = 0.0003; P−/- versus PVUS/VUS_02, P = 0.0037). (B) Fibroblasts from an individual with novel biallelic missense variants in AP4B1 (PVUS/VUS_01), an asymptomatic individual with a heterozygous variant in AP4B1 (P+/-_01) and an AP-4-HSP patient with biallelic loss-of-function variants in AP4B1 (P−/−_01) stained with phalloidin (actin marker) and antibodies against TGN46 (trans-Golgi network marker) and ATG9A. Right panel shows magnification of inserts. Dotted line indicates from (B). (C and D) Western blot of whole-cell lysates from fibroblasts of PVUS/VUS_01 compared to P−/−_01 and two AP-4-HSP patients with biallelic loss-of-function variants in AP4B1 (P−/+_01 and P−/-_07). Fibroblasts from PVUS/VUS_01 show levels of AP4B1, AP4E1 and ATG9A that are not different compared to P−/−_01, whereas levels of AP4B1 are robustly reduced and levels of ATG9A elevated in P−/-_01 and P−/-_07 (one-way ANOVA with Dunnett’s post hoc test, P < 0.0001). Bar graph shows the mean and standard deviation of 4–6 (PVUS/VUS_01) and 5 (P−/+_01 and P−/-_07) samples. (E and F) Western blot of whole-cell lysates from fibroblasts of PVUS/VUS_02 compared to P−/+_01 and two AP-4-HSP patients with biallelic loss-of-function variants in AP4B1 (P−/+_01 and P−/-_07). Fibroblasts from PVUS/VUS_02 show a reduction of AP4B1 and AP4E1, indicative of lower AP-4 complex formation. There is, however, no significant change in ATG9A compared to P−/-_01 (one-way ANOVA with Dunnett’s post hoc test, P < 0.0001). Bar graph shows the mean and standard deviation of 6 (PVUS/VUS_02) and 4 (P−/+_01 and P−/-_07) samples. (G) Co-immunoprecipitation of AP4E1 and AP4B1 demonstrates reduced binding of both subunits in lysates from PVUS/VUS_02 compared to a control (P−/+_04). For comparison, the same experiment with lysates from a patient with biallelic loss-of-function variants in AP4M1 is shown (P−/-_11), demonstrating near complete absence of AP-4 complex assembly. (H and I) Fibroblasts from an individual with novel biallelic missense variants in AP4B1 (PVUS/VUS_02), an asymptomatic control (P−/+_04) and an AP-4-HSP patient with biallelic loss-of-function variants in AP4M1 (P−/-_11) stained with antibodies against AP4E1 and TGN46 showing reduced levels of AP4E1 associated with the TGN in PVUS/VUS_02. No significant change in ATG9A localization is found. ***P < 0.001; ****P < 0.0001.
**Nucleofection**

AP4B1 plasmid DNA constructs were purchased from Twist Bioscience (Supplementary File 2). Plasmid to express pmaxGFP was purchased from Lonza (Lonza, #VCA-1003). Nucleofection of AP4B1KO SH-SY5Y cells with full-length wild type AP4B1 or AP4B1 carrying variants found in PVUS/VUS_01 and PVUS/VUS_02 was carried out using the Cell Line Nucleofector Kit V (Lonza, #VCA-1003) and Amaza Nucleofector 2b device (Lonza, #AAB-1001). Per 2 × 10^6 cells, 2 μg of AP4B1 plasmid and 2 μg of GFP plasmid were used. After nucleofection cells were resuspended in differentiation medium with retinoic acid and seeded onto 96-well plates (Greiner Bio-One, Cat# 655090) at a density of 14 × 10^3 per well. After 48 h, cells were fixed using 4% paraformaldehyde (PFA) and stained and imaged as described below.

**Immunocytochemistry**

Automation of each step in the immunocytochemistry protocol was maximized by using automated pipettes and reagent dispensers (Thermo Fisher Scientific Multidrop Combi Reagent Dispenser, Integra VIAFLO 96/384 liquid handler, Integra VOYAGER pipette). After 24 h in a humidity-controlled incubator, plates were fixed using 3% PFA, permeabilized with 0.1% saponin in phosphate-buffered saline (PBS) and blocked in PBS containing 1% bovine serum albumin (BSA) and 0.01% saponin (blocking solution). Primary antibody (diluted in blocking solution) was added for 1 h at room temperature. Plates were washed three times in blocking solution and then fluorochrome-conjugated secondary antibody, Hoechst 33258 and Phalloidin were added for 1 h at room temperature. Plates were washed three times with PBS. For experiments shown in Fig. 5H and I, cells were grown onto 13 mm glass coverslips and fixed in 3% formaldehyde in PBS or, for AP4E1 labelling, ice-cold methanol. Formaldehyde fixed cells were permeabilized and blocked in PBS with 0.5% BSA and 0.01% saponin. Methanol fixed cells were blocked in 0.5% BSA. Primary antibody (diluted in blocking solution) was added for 60 min at room temperature. Coverslips were washed three times in PBS and then fluorochrome-conjugated secondary antibody was added in blocking solution for 40 min at room temperature. Coverslips were then washed three times in PBS. Widefield and confocal images were captured on a Zeiss LSM 710 confocal microscope on an inverted Axiolmager Z1 using a Zeiss Plan Apochromat 63× oil immersion objective (NA 1.4) and ZEN Black software version 2.3. Images were always processed in the same way with the same exposure times and the same manipulations to optimize brightness and contrast. For experiments shown in Fig. 6, plates with differentiated SH-SY5Y cells were fixed in 4% PFA at 48 h after nucleofection, permeabilized with 0.1% saponin in PBS and blocked in 1% BSA/0.01% saponin in PBS. Primary antibody diluted in blocking solution was added for 1 h at room temperature. Plates were washed three times in blocking solution and fluorochrome-conjugated secondary antibodies and Hoechst 33258 were added for 30 min at room temperature.

**High-content imaging and automated image analysis**

High-throughput confocal imaging was performed on an ImageXpress Micro Confocal Screening System (Molecular Devices) using an experimental pipeline modified from the pipeline described in Behne et al.11 For experiments shown in Figs 3–5A and B, images were acquired using a 20× S Plan Fluor objective (NA 0.45 μM, WD 8.2–6.9 mm). Per well, 9 fields were acquired in a 3 × 3 format (96-well plates) or 4 fields in a 2 × 2 format (384-well plates). Each experiment was performed in biological duplicate on two separate plates with each plate carrying loss-of-function (LoF) and matched wild-type (WT)/LoF cells. Analysis was performed using a customized image analysis pipeline in MetaXpress (Molecular Devices): Briefly, cells were identified based on the presence of DAPI signal inside a phalloidin-positive cell body. Sequential masks were generated for (1) the TGN by outlining the area covered by TGN marker TGN46 (TGN46+ area) and (2) for the cell area outside the TGN (actin+ area minus TGN46+ area). ATG9A fluorescence intensity was measured in both compartments in each cell and the ‘ATG9A’ ratio was calculated by dividing the ATG9A fluorescence intensity in the TGN by the ATG9A fluorescence intensity in the remaining cell body. For experiments shown in Fig. 6, images were acquired using a 40× S Plan Fluor objective (NA 0.60 μM, WD 3.6–2.8 mm). Per well, 64 fields were acquired in an 8×8 format. Automated image analysis was performed as described above using the eGFP signal, instead of phalloidin, to outline the cytoplasm. Z’-factor robust values (Z’ robust = 1–3*[mad(a)+mad(b)]/abs[median(a)–median(b)]) and strictly standardized mean difference (SSMD) were calculated for each plate and only plates that met the predefined quality metrics of a Z’-factor robust ≥ 0.3 and SSMD ≥ 3 were included in subsequent analyses.

**Western blot**

Western blotting was done as previously described.36 Western blots shown in Fig. 5G were developed using chemiluminescence detection of HRP-conjugated secondary antibody using Amersham ECL Prime Western Blotting Detection Reagent (GE Healthcare) and X-ray film.

**Immunoprecipitation**

For immunoprecipitations of AP4E1 and AP4B1, equal starting material (similar passage number and growth rate) of fibroblasts from P+/−_4 (unaffected control), P+/−_02 (patient with novel biallelic missense variants in AP4B1) and P−/−_11 (patient with biallelic loss-of-function variants in AP4M1) were lysed in PBS with...
recovered in NuPAGE LDS Sample Buffer at 75 X-100, once in PBS, and the immunoprecipitates were added, and samples were incubated for a further 45 min. A sepharose beads (GE Healthcare, #17-0780-01) were axis limits). Data shown in Fig. 4F were compared using indicating the median (334 data points were outside the sh o w s v i o l i n p l o t s o f a l l d a t a p o i n t s w i t h h o r i z o n t a l l i n e s

Statistical analysis

Statistical analysis was performed in GraphPad Prism version 9.0. Normality of the data was assessed with the Shapiro–Wilk test and QQ plots. Figures 3C, 4B, D–F, 5A, D and F show the mean with standard deviation. Figure 4A shows violin plots of all data points with horizontal lines indicating the median (334 data points were outside the axis limits). Data shown in Fig. 4F were compared using the Mann–Whitney test. For experiments shown in Fig. 4B, D and E groups were compared using Kruskal–Wallis test with post hoc Dunn’s test for multiple comparisons. Data shown in Figs 5A, D, F and 6B were compared using one-way ANOVA with Dunnett’s post hoc test after a normal distribution was established. The Receiver Operator Characteristic (ROC) curve shown in Fig. 4C was computed by pooling all values for patients (PVT_1 – PVT_18) and heterozygous controls (PVT_1 – PVT_17). For Figs 3D and 4G, a simple linear regression analysis was performed and the Pearson correlation coefficient was calculated. Statistical significance was determined at P < 0.05 for all analyses.

Data availability

Data are available from the corresponding authors upon reasonable request. Fibroblast lines generated in this study are available with a material transfer agreement.

Results

Clinical and molecular characterization

We established fibroblast lines from 20 individuals with biallelic variants in subunits of the AP-4 complex
### Table 1: Clinical and molecular characterization of the study cohort

| Patient ID | AP4-HSP subunit | Allele 1 | Allele 2 | CADD PHRED score | ACMG classification a | Sex | Age | SPRS | 4-FMS | AP-4-HSP core features | ATG9A ratio |
|------------|----------------|----------|----------|------------------|-----------------------|-----|-----|------|-------|-------------------------|------------|
| P/−_01     | AP4B1          | c.1345A>T (p.Arg449Ter) (Nonsense) | c.1160_1161delCA (p.Thr387ArgfsTer30) (Frameshift) | 36/33              | Pathogenic/pathogenic | M   | 3 y 11 m | 18   | 2     | 9/9                     | 1.65       |
| P/−_02     | AP4B1          | c.530_531insA (p.Asn178GlufsTer20) (Frameshift) | c.11-4-2A>C p. (Splice site) | 32/34              | Pathogenic/pathogenic | F   | 5 y 3 m | 15   | 3     | 7/9                     | 1.59       |
| P/−_03     | AP4B1          | c.1216C>T (p.Arg406Ter) (Nonsense) | c.1328T>C (p.Leu443Pro) (Nonsense) | 41/28.1            | Pathogenic/uncertain | F   | 4 y 11 m | 35   | 4     | 8/9                     | 1.44       |
| P/−_04     | AP4B1          | c.1216C>T (p.Arg406Ter) (Nonsense) | c.1160_1161delCA (p.Thr387ArgfsTer30) (Frameshift) | 41/28.1            | Pathogenic/uncertain | F   | 4 y 10 m | 22   | 3     | 8/9                     | 1.70       |
| P/−_05     | AP4B1          | c.11-4-2A>C p. (Splice site) | c.11-4-2A>C p. (Splice site) | 34/34              | Pathogenic/pathogenic | F   | 7 y 5 m | 32   | 3     | 8/9                     | 1.71       |
| P/−_06     | AP4B1          | c.1160_1161delCA (p.Thr387ArgfsTer30) (Frameshift) | c.1160_1161delCA (p.Thr387ArgfsTer30) (Frameshift) | 33/33              | Pathogenic/pathogenic | M   | 4 y 8 m | 22   | 3     | 8/9                     | 1.39       |
| P/−_07     | AP4B1          | c.644delG (p.Leu222GlufsTer31) (Frameshift) | c.1177C>T (p.Arg393Ter) (Nonsense) | 28.9/40            | Pathogenic/pathogenic | M   | 1 y 9 m | NA   | 4     | 8/9                     | 1.75       |
| P/−_09     | AP4B1          | c.1608_1609insGA (p.Lys537GlnfsTer18) (Frameshift) | c.1608_1609insCA (p.Lys537GlnfsTer18) (Frameshift) | 32/32              | Pathogenic/pathogenic | M   | 7 y 2 m | 22   | 2     | 8/9                     | 1.63       |
| P/−_10     | AP4M1          | c.910C>T (p.Arg306Ter) (Nonsense) | c.910C>T (p.Arg306Ter) (Nonsense) | 39/39              | Pathogenic/pathogenic | F   | 31 y 1 m | NA   | 4     | 8/9                     | 1.36       |
| P/−_11     | AP4M1          | c.218dupA (p.Asn73GlufsTer43) (Frameshift) | c.851A>C (p.Tyr284Ser) (Nonsense) | 24.5/29.5         | Pathogenic/uncertain | M   | 9 y 1 m | 41   | 4     | 9/9                     | 1.50       |
| P/−_12     | AP4M1          | c.910C>T (p.Arg306Ter) (Nonsense) | c.694dupG (p.Glu232GlufsTer21) (Frameshift) | 39/33              | Pathogenic/pathogenic | M   | 18 m 29 | 4    | 9/9   | 1.46                    |           |
| P/−_13     | AP4M1          | c.1025 + 1dup T p. (Splice site) | c.205A>G (p.Glu62ArgfsTer21) (Frameshift) | 39/33              | Pathogenic/pathogenic | M   | 5 y 10 m | 0    | 2     | 6/9                     | 1.37       |
| P/−_14     | AP4S1          | c.138 + 3_138 + 6delAAAT p. (Splice site) | c.138 + 3_138 + 6delAAAT p. (Splice site) | 34/22.4            | Uncertain significance/uncertain significance | M   | 4 y 11 m | NA   | 3     | 7/9                     | 1.39       |
| P/−_15     | AP4S1          | c.284 + 1G>T p. (Splice site) | c.284 + 1G>T p. (Splice site) | 35/35              | Pathogenic/pathogenic | M   | 3 y 6 m | 19   | 3     | 9/9                     | 1.52       |
| P/−_16     | AP4S1          | c.289C>T (p.Arg97Ter) (Nonsense) | c.289C>T (p.Arg97Ter) (Nonsense) | 50/50              | Pathogenic/pathogenic | F   | 4 y 2 m | 22   | 2     | 9/9                     | 1.64       |
| P/−_17     | AP4S1          | c.49dupT (p.Ser17GlufsTer21) (Frameshift) | c.49dupT (p.Ser17GlufsTer21) (Frameshift) | 30/30              | Pathogenic/pathogenic | F   | 12 y 9 m | NA   | 3     | 8/9                     | 1.51       |
| P/−_18     | AP4S1          | c.49dupT (p.Ser17GlufsTer21) (Frameshift) | c.238_239insG (p.Leu80SerfsTer3) (Frameshift) | 30/33              | Pathogenic/pathogenic | M   | 17 y 1 m | 24   | 3     | 8/9                     | 1.56       |
| P/PLUS/PLUS_01 | AP4B1        | c.1024G>T (p.Asp342Ter) (Missense) | c.868C>T (p.Arg290Cys) (Missense) | 28.6/25.5         | Uncertain significance/uncertain significance | F   | 3 y 0 m | NA   | 4     | 3/9                     | 1.16       |
| P/PLUS/PLUS_02 | AP4B1        | c.409A>G (p.Arg137Gly) (Missense) | c.409A>G (p.Arg137Gly) (Missense) | 24.4/24.4         | Uncertain significance/uncertain significance | F   | 3 y 1 m | NA   | 2     | 1/8                     | 1.27       |

F, female; M, male; m, months; NA, not available; SPRS, Spastic Paraplegia Rating Scale; y, years. Reference sequences: AP4B1: NM_004722.4; AP4M1: NM_007347.5; AP4S1: NM_007077.4.

aClassification of variants based on ACMG criteria using VarSome (Kopanos et al., 2019). CADD PHRED scores were computed using version 1.6 (https://cadd.gs.washington.edu/score).27
(Table 1, Supplementary Table 1), as well as 14 fibroblast lines from parents or siblings (confirmed asymptomatic, heterozygous carriers) and 10 healthy, unrelated control individuals (Supplementary Table 2). Demographic, clinical and genetic information are provided in Table 1. Clinical information of 13 patients has been reported previously. In the current cohort, 18/20 of referred individuals showed ≥6 of the 9 core clinical and radiographic features of AP-4-HSP (according to Ebrahimi-Fakhari et al., a list can be found in Supplementary Table 3). The majority of patients presented with developmental delay and subsequent intellectual disability (19/20), progressive spasticity (19/20), microcephaly (17/20) and epilepsy (14/20). Most individuals ambulated with assistance or used a wheelchair as indicated by 4-Stage Mobility Scores of III (8/20) and IV (7/20). The mean Spastic Paraplegia Rating Scale (SPRS) score was 24.4 ± 10.8 (SD) indicating a moderate degree of corticospinal tract dysfunction and associated complications. Common neuroimaging features consisted of a thin corpus callosum (17/20), abnormal signal in the periventricular white matter (18/20) and ex-vacuo ventriculomegaly (15/20).

**AP4B1** was most commonly mutated in our cohort. Variants covered a wide spectrum including frameshift/nonsense (22/40 alleles), missense (10/40 alleles) and splice site variants (8/40 alleles), in homozygous or compound heterozygous states (Table 1). Eight alleles carried variants of uncertain significance by ACMG criteria, including one individual with a compound heterozygous splice site and missense variant in AP4S1 (P.13, and two individuals with a compound heterozygous or homozygous missense variant in AP4B1, respectively (PVUS/VUS_01 and PVUS/VUS_02). CADD scores for all variants were above the recommended cut-off for significance of 20.

Individuals PVUS/VUS_01 and PVUS/VUS_02 presented with atypical clinical features and novel variants in AP4B1 (Table 1, Supplementary Table 1), and were thus referred for a functional interpretation of their variants.

Briefly, PVUS/VUS_01 is a 3-year-old white female, born to non-consanguineous parents of mixed-European heritage, who presented in the neonatal period with refractory epilepsy with myoclonic seizures, followed by severe global developmental delay and hypotonia. Brain MR imaging was normal. Exome sequencing showed compound-heterozygous missense variants in AP4B1 (NM_001253852.3), classified as a variant of unclear significance (VUS) and a likely benign variant, respectively. The c.1024G>T (p.Asp342Tyr) variant showed an allele frequency of 0.0000398 in gnomAD exomes, with a homozygous allele count of 0. There were 7 benign predictions from BayesDel_addAF, DEOGEN2, EIGEN, MVP, MutationAssessor, PrimateAI and REVEL versus 6 pathogenic predictions from DANN, FATHMM-MKL, LIST-S2, M-CAP, MutationTaster and SIFT. CADD PHED scores (version 1.6) were 28.6 and 25.5, respectively (Table 1).

PVUS/VUS_02 is a 41-year-old white woman of Belgian ancestry, with no history of parental consanguinity, who presented with adult-onset mild spastic paraplegia, mild intellectual disability, ataxia and dysarthria. There was no history of seizures or microcephaly. She continues to ambulate with a spastic gait and is independent with most activities of daily living. Brain MR imaging showed mild cerebellar atrophy but a normally formed corpus callosum, lateral ventricles and periventricular white matter. Exome sequencing showed a homozygous missense variant in AP4B1 (NM_001253852.3), classified as a variant of uncertain significance. The variant, c.409A>G (p.Arg137Gly), was absent from gnomAD exomes and genomes and was predicted to be pathogenic based on 11 pathogenic predictions from BayesDel_addAF, DANN, DEOGEN2, EIGEN, FATHMM-MKL, LIST-S2, M-CAP, MutationAssessor, MutationTaster, REVEL and SIFT, versus 2 benign predictions from MVP and PrimateAI. The CADD score was 24.4.

### AP-4 subunit structures with variant distribution and computational scores

The distribution of variants in the AP-4 subunit genes of individuals included in the AP-4-HSP International Registry and Natural History Study (ClinicalTrials.gov Identifier: NCT04712812) is shown in Figs 1 and 2. The CADD score of each variant is indicated and a model of CADD scores is provided for all possible missense variants across the protein secondary structures of AP4B1, AP4M1, AP4E1 and AP4S1 (similar analyses for M-CAP and REVEL scores can be found in Supplementary Fig. 1, data are summarized in Supplementary File 1). All types of variants observed in patients were predominately located in the known functional domains. This observation was particularly evident for missense variants (blue dots) in AP4B1 and AP4M1, which were exclusively located in either the Adaptin N terminal region (in the case of AP4B1, Fig. 1A) or the Adaptor complexes medium subunit domain (in the case of AP4M1, Fig. 1B). Novel missense variants present in PVUS/VUS_01 and PVUS/VUS_02 localized to the Adaptin N terminal region of AP4B1. In AP4B1, a few truncating or splice variants were located in the region between the functional domains (Fig. 1A). However, the C-terminal B2-adapt-app_C domain of AP4B1 is notable, since no disease-associated variants have been identified here to date. In AP4B1, AP4E1 and AP4S1, higher CADD scores in
functional domains indicated greater restrain for missense variation (Figs 1A, 2A and B). AP4M1 deviates from this pattern since no lowering of CADD scores was found for missense variants that fall between the functional domains (Fig. 1B). The observed higher conservation of this shortest linker region of all AP-4 proteins possibly indicates that this region is important to precise inter-domain spacing and positioning.

Validation of the ‘ATG9A ratio’ as an indicator of AP-4 function and a diagnostic marker for AP-4-HSP

The AP-4 complex mediates transport of transmembrane proteins from the TGN to the periphery of the cell. Three independent groups identified the core autophagy protein ATG9A as a major cargo of AP-4. In a small cohort of AP-4-HSP patients, using conventional immunocytochemistry, we previously showed that ATG9A accumulates in the TGN.

Building on this finding, we developed an automated, high-throughput imaging platform that allows us to measure the intracellular distribution of ATG9A in thousands of patient-derived fibroblasts in a single run (Fig. 3A and B). Specifically, we determined the ratio of ATG9A fluorescence signal in the area of the TGN in relation to the signal in the remainder of the cell (Fig. 3B). Using this ‘ATG9A ratio’, we established that our assay has favourable metrics for cell-based high-throughput assays (Z-factor robust > 0.3, SSMD > 3; Fig. 3C and Supplementary Fig. 2) and reproducibility between replicates (Fig. 3D). Confirming that our assay provides a surrogate of AP-4 function and is sensitive to its restoration, we found that re-expression of the missing AP4B1 subunit in fibroblasts with biallelic truncating variants in AP4B1 restored the ATG9A ratio to a level close to controls (Fig. 3E).

We found that fibroblast lines from patients with typical clinical features of AP-4-HSP (≥6 core clinical features), most (16/18) with at least one loss-of-function variant (frameshift, nonsense or essential splice site), showed a significant increase in the ATG9A ratio compared to heterozygous and healthy unrelated controls (Fig. 4A and B, Table 2). This increase was ~1.3-fold compared to heterozygous controls from the same assay plate (Table 2). Figure 4A shows violin plots demonstrating the distribution of the ATG9A ratio on the level of individual cells with a mean of 6765 ± 4536 (SD) cells per cell line. Figure 4B shows a grouped analysis summarized by assay plate with two biological replicates per cell line. ROC analysis showed that the ATG9A ratio can separate patients with biallelic loss-of-function variants from heterozygous controls with robust diagnostic power (area under the curve 0.85, 95% confidence interval 0.849–0.852) (Fig. 4C). Individuals with AP4M1-associated SPG50 showed a lower mean ATG9A ratio compared to individuals with AP4B1-associated SPG47 (Fig. 4D).

Fibroblasts were derived from three patients with compound heterozygous missense variants where one allele was classified as of uncertain significance and the other as pathogenic (P<–3, P<–4, P<–11) showed an ATG9A ratio that was not significantly different from that of patients with two variants classified as pathogenic (Fig. 4E). This finding supports the pathogenicity of the variants. Similarly, in one individual with a novel missense variant in trans with a novel splice-site variant (P<–13), both previously classified as of uncertain significance, we found an ATG9A ratio that confirms the detrimental impact of these variants on AP-4 function (Fig. 4E). In a grouped analysis, patients with one or more missense alleles tended to have lower ATG9A ratios than patients with two nonsense, frameshift, or splice site variants, though this difference did not reach statistical significance (Fig. 4F). Exploring correlations with in silico prediction tools, we found that there was a trend, but no significant correlation, between the CADD scores of each patient’s variants and their corresponding ATG9A ratio levels (Fig. 4G).

The ATG9A ratio can provide a functional assessment of novel missense variants

Having established the ATG9A ratio in a well-characterized cohort of patients with AP-4-HSP, we next explored if our assay could be used to functionally characterize novel variants in AP-4 subunit genes. We found that fibroblasts from two individuals with biallelic missense variants in AP4B1 and atypical phenotypes in AP4B1 and AP4E1, which can be used as a surrogate for AP-4 stability, were not different compared to controls (Fig. 5C and D). In addition, the level of ATG9A was unaffected, whereas ATG9A levels were increased in cells from patients with bona fide AP-4-HSP (Fig. 5C and D). Since these proteins are known to change with the loss of AP-4 function, including in AP-4 knockout mice and cells from AP-4-HSP patients, the

| Table 2 ATG9A ratio can discriminate AP-4-HSP patients from controls |
|---------------------------------------------------------------|
| **Typical AP-4-HSP patients (n = 18)** | **Heterozygous carriers (n = 14)** |
| ATG9A ratio | ATG9A ratio |
| 1.54 ± 0.13 (mean, SD), 95% CI of mean: 1.50–1.58 | 1.21 ± 0.05 (mean, SD), 95% CI of mean: 1.18–1.23 |
| 1.54 (median), 95% CI of median: 1.49–1.62 | 1.20 (median), 95% CI of median: 1.17–1.24 |
absence of change argues against a functionally-relevant impact of the novel missense variants in \( P_{VUS/VUS}^1 \).

In \( P_{VUS/VUS}^2 \), as in \( P_{VUS/VUS}^1 \), the ATG9A ratio was significantly lower than that of patients with typical AP-4-HSP and biallelic loss-of-function variants (Fig. 5A). However, looking at protein levels in whole-cell lysates, we found a reduction of AP4B1 and AP4E1, by about 30% compared to heterozygous controls, although no change in ATG9A protein levels (Fig. 5E and F). To further investigate the effect of the variant on AP-4, we used immunocytochemistry experiments (Fig. 5H and I). Immunostaining revealed a reduced presence of AP4E1 at the TGN, suggestive of reduced levels of AP-4 complex associated with membranes (Fig. 5H). In contrast, there was no significant impact on ATG9A at the TGN (Fig. 5I). Taken together, these findings argue that the homozygous c.409A>G (p.Arg137Gly) missense variant leads to reduced AP-4 complex formation. Supporting this hypothesis, immunoprecipitation experiments showed reduced binding of AP4E1 and AP4B1 (Fig. 5G). Importantly, however, there was residual complex formation, whereas no apparent complex formation was seen in cells with biallelic loss of function variants (Fig. 5G and I).

Collectively, findings in \( P_{VUS/VUS}^2 \) indicate that while this homozygous missense variant leads to lower AP-4 complex formation, the level of AP-4 complex expression appears to be sufficient for efficient sorting of ATG9A at the TGN in fibroblasts.

To further characterize the functional impact of each variant present in \( P_{VUS/VUS}^1 \) and \( P_{VUS/VUS}^2 \), we expressed each mutant allele in human AP4B1 knockout SH-SY5Y cells, a neuronal model of AP-4 deficiency. AP4B1 knockout SH-SY5Y cells show an elevated ATG9A ratio, indicating accumulation of ATG9A in the TGN area (Fig. 6A and B). Using our high-throughput imaging platform, we established that this assay can reliably detect differences between negative (AP4B1 knockout) and positive (AP4B1 wild type) controls (mean: \( Z^* \)-factor robust: 0.92, SSMD: 30.9). We found that re-expression of each mutant allele (c.1024G>T, c.868C>T, c.409A>G) over 48 h led to a significant reduction in the ATG9A ratio, providing evidence that they, at least in part, restore AP-4 function in a neuronal model of AP-4 deficiency.

In summary, based on data in well-characterized AP-4-HSP patients and controls, we establish that an elevated ATG9A ratio in patient-derived fibroblasts indicates clinically relevant AP-4 dysfunction with a high degree of sensitivity and specificity, providing a diagnostic functional assay for cases with atypical features and variants of uncertain significance.

Discussion

In the absence of a biochemical biomarker in blood, urine or CSF, a functional assay in patient-derived fibroblasts can aid the interpretation of novel gene variants and help establish a clinical diagnosis. Having a functional assay is of particular importance to rare disorders with a wide or non-specific clinical spectrum. Cell-based functional assays similar to ours exist for a small number of disorders, mainly inborn errors of metabolism, where the function of an enzyme or accumulation of a metabolite can be assayed. Such disorders include other forms of HSP, for example, SPG5 where oxysterol levels can be measured \(^{42}\) and SPG26 where GM2 synthase activity can be approximated, \(^{43}\) and lysosomal storage diseases, for example, the use of filipin staining used in the diagnosis of Niemann–Pick disease type C. \(^{44}\)

With just over 200 individuals identified to date, \(^3\) AP-4-HSP is an ultra-rare, yet paradigmatic, form of childhood-onset HSP and a genetic mimic of cerebral palsy. The diagnosis relies on molecular testing, and there is often a significant diagnostic delay. \(^{45}\) To aid the diagnostic process and to provide a tool for gene variant interpretation, we developed a diagnostic functional assay that measures a known and well-established downstream target of AP-4.

ATG9A is a transmembrane autophagy-related protein and cycles between the TGN and ATG9A vesicles. There is increasing evidence that ATG9A acts as a lipid scramblase that enables the expansion of nascent autophagosome membranes. \(^{15–17}\) In neurons, autophagosomes form in the distal axon \(^{46,47}\) and are subject to active transport. \(^{36,48–50}\) Thus, transport of ATG9A to distal axons may act as a cue to compartmentalize autophagosome biogenesis. \(^{39,41,51,52}\) From recent work, \(^{9,11,36,41,53}\) and overlapping phenotypes of AP-4 \(^{39,41,54}\) and Atg9a knockout mice, the following working model for AP-4 deficiency emerges: (i) AP-4 is required for sorting of ATG9A from the TGN; (ii) loss-of-function variants in any of the four AP-4 subunits lead to a loss of AP-4 assembly and function; (iii) ATG9A accumulates in the TGN leading to a reduction of axonal delivery of ATG9A; and (iv) lack of ATG9A at the distal axon impairs autophagosome biogenesis and axonal function, leading to the length-dependent axonal degeneration. While loss of AP-4 function clinically leads to a neurological disorder, the mislocalization of ATG9A can be assayed in multiple cell types and tissues, including in fibroblasts, as shown here.

Our assay employs automated high-throughput microscopy of ATG9A in patient-derived fibroblasts and can reliably detect reduced AP-4 function and thus AP-4-associated HSP. The assay can be performed following a standard diagnostic skin biopsy and using standard laboratory resources. Following fibroblast derivation, results can be obtained within days. Based on the data from 18 patients with genetically and clinically confirmed AP-4-HSP, we define a value between 1.49–1.62 (representing the 95% CI of the median in this cohort) as strongly indicative of clinically significant AP-4 dysfunction. ROC analysis shows an AUC of 0.85 indicating robust separation of patients and controls, with an ATG9A
ratio of >1.37 showing a sensitivity of 69% and a specificity of 89% for detecting AP-4-HSP. Relying on a ratio rather than an absolute fluorescence value and by deriving data from thousands of cells in each run, our assay performs robustly and reliably. Controls are included on each assay plate and suitable positive controls may include fibroblast lines from patients with biallelic early frameshift variants and established loss of function, whereas negative controls can be fibroblasts derived from a family member with a heterozygous variant or unrelated wild type controls.

In cases where the pathogenicity of variants is unclear and/or the phenotype deviates from core clinical features present in the majority of AP-4-HSP patients, our platform can provide an approximation of AP-4 function. As demonstrated in two atypical cases with novel missense variants, this can help substantiate or disprove a clinical diagnosis and, thus, can aid the diagnostic workup, with implications for counselling and treatment. Findings in $P^{VUS/VUS}_02$ illustrate that biallelic missense variants may lower the probability of AP-4 complex formation, yet this might not lead to a lack of trafficking of AP-4 cargo. There is likely a threshold of residual expression that allows sufficient AP-4 complex to form and cargo to be transported. This notion is supported by additional re-expression experiments in $AP4B1^KO$ SH-SY5Y cells, a human neuronal model for AP-4 deficiency. We found that expression of the c.409A>G allele reduced the ATG9A ratio 48 h after transfection, yet to a lower degree compared to expression of the full-length wild-type protein or the c.1024G>T and c.868C>T alleles found in $P^{VUS/VUS}_01$. Thus, it remains possible that the c.409A>G variant conveys a partial reduction in AP-4 function possibly by reducing the amount of AP-4 assembly. From knockdown experiments of AP-1, however, we estimate that expression of adaptor protein complexes would have to be reduced substantially, i.e. >90%, to lead to a significant impact on downstream targets.55,56

We recognize that a subtle impairment of ATG9A trafficking in neurons, particularly neuronal populations known to be vulnerable to changes in autophagy57,58 and involved in AP-4-HSP,59 might not be reflected in fibroblasts or other cell culture-based experiments. We also acknowledge that our assay relies on ATG9A distribution only and does not take into account other cargo proteins that are potentially important in neurons.60,61 Thus, an interpretation of results from our fibroblast-based assay should integrate detailed clinical information, neuroimaging, molecular testing and in silico predictions, including those illustrated in Figs 1 and 2 and Supplementary Fig. 1.

The overlap of CADD PHRED scores of missense variants across $AP4B1$, $AP4E1$ and $AP4S1$ with the annotated domains underscores their relevance for function and variant interpretation (i.e. ACMG criterion PM1), but as shown for variants in individuals $P^{VUS/VUS}_01$ and $P^{VUS/VUS}_02$ this has limitations. In a large cohort, our functional assay used in combination with variant location and in silico prediction scores may provide an opportunity to further refine the impact of missense variants, aid the interpretation and classification of variants and enable further exploration of differences between subunits and functional domains in combination with structural biology approaches.61 This could be done post hoc focusing on missense variants identified in individuals through molecular testing or prospectively using a massive parallel screen of all possible missense variants, as has been recently shown for a cancer predisposition gene.62

For the future development of biomarkers, it will be important to establish if changes in ATG9A can be detected in other specimens including in blood, saliva or cerebrospinal fluid, which would allow for the investigation of longitudinal changes with disease progression or the impact of interventions.

In conclusion, we here establish a high-throughput ATG9A translocation assay in patient-derived fibroblasts as a diagnostic test for AP-4-HSP. Our approach enables the functional validation of novel variants in $AP4B1$, $AP4E1$, $AP4S1$ and $AP4E1$ and can thus provide additional support for a clinical and molecular diagnosis of this debilitating childhood neurological disorder.

Supplementary material
Supplementary material is available at Brain Communications online.

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Competing interests

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