Hereditary spastic paraplegia is a novel phenotype for GJA12/GJC2 mutations

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Recessive mutations in GJA12/GJC2, the gene that encodes the gap junction protein connexin47 (Cx47), cause Pelizaeus-Merzbacher-like disease (PMLD), an early onset dysmyelinating disorder of the CNS, characterized by nystagmus, psychomotor delay, progressive spasticity and cerebellar signs. Here we describe three patients from one family with a novel recessively inherited mutation, 99C>G (predicted to cause an Ile→Met amino acid substitution; I33M) that causes a milder phenotype. All three had a late-onset, slowly progressive, complicated spastic paraplegia, with normal or near-normal psychomotor development, preserved walking capability through adulthood, and no nystagmus. MRI and MR spectroscopy imaging were consistent with a hypomyelinating leukoencephalopathy. The mutant protein forms gap junction plaques at cell borders similar to wild-type (WT) Cx47 in transfected cells, but fails to form functional homotypic channels in scrape-loading and dual whole-cell patch clamp assays. I33M forms overlapping gap junction plaques and functional channels with Cx43, however, I33M/Cx43 channels open only when a large voltage difference is applied to paired cells. These channels probably do not function under physiological conditions, suggesting that Cx47/Cx43 channels between astrocytes and oligodendrocytes are disrupted, similar to the loss-of-function endoplasmic reticulum-retained Cx47 mutants that cause PMLD. Thus, GJA12/GJC2 mutations can result in a milder phenotype than previously appreciated, but whether I33M retains a function of Cx47 not directly related to forming functional gap junction channels is not known.

Keywords: spastic paraplegias; Pelizaeus-Merzbacher-like disease; gap junction; connexin; oligodendrocyte

Abbreviations: Cx47 = connexin47; DTRs = deep tendon reflexes; ER = endoplasmic reticulum; HSP = hereditary spastic paraplegia; PMD = Pelizaeus-Merzbacher disease; PMLD = Pelizaeus-Merzbacher-like disease
Introduction

Pelizaeus-Merzbacher disease (PMD) is an X-linked disorder caused by mutations in PLP1, the gene encoding proteolipid protein, the main protein in CNS myelin. Classic PMD affects boys and is characterized by nystagmus and impaired psychomotor development within the first year of life, followed by progressive spasticity, ataxia, choreoathetosis and diffuse white matter changes on MRI (Nave and Boespflug-Tanguy, 1996; Hudson et al., 2004; Inoue, 2005). PLP1 mutations may also cause a more severe 'connatal' PMD phenotype, which is characterized by dramatic psychomotor impairment from birth, with hypotonia, quadriplegia, nystagmus, seizures and early death. Yet other PLP1 mutations cause a much milder disease, a 'pure' spastic paraplegia type 2 (SPG2), or a more 'complicated' form, with onset in childhood or adolescence, and dysarthria, mental retardation and ataxia (Garbern et al., 1999; Hudson et al., 2004; Garbern, 2007).

Pelizaeus-Merzbacher-like disease (PMLD) is clinically and neuroradiologically similar to classic PMD, but is not associated with PLP1 mutations. Recessive mutations in GJA12 are one cause of PMLD (Uhlenberg et al., 2004; Bugiani et al., 2006; Salvati et al., 2007; Woll et al., 2007; Henneke et al., 2008). GJA12/GJC2 (GJA12 was recently renamed GJC2; http://www.genenames.org/genefamily/gj.php) encodes connexin47 (Cx47), which is a member of the connexin family, highly conserved integral membrane proteins usually named according to their predicted molecular mass (Willecke et al., 2002). Connexins form gap junctions, which are intercellular channels that form between apposed cell membranes to permit the diffusion of ions and small molecules typically less than 1000 Da (Bruzzone et al., 1996). Six connexins oligomerize into a hemichannel (or connexon), and two apposing hemichannels form the gap junction; aggregates of tens to thousands of intercellular channels form a gap junction plaque. The potential diversity of gap junction composition is immense, as over 20 mammalian connexins have been described. Hemichannels may be homomeric, containing one type of connexin, or heteromeric, containing more than one type. Gap junctions are termed homotypic if the apposed hemichannels contain the same connexin, and heterotypic if they contain different connexins (Kumar and Gilula, 1996).

Anatomical and functional studies of the mammalian CNS have demonstrated that astrocytes and oligodendrocytes are coupled by gap junctions, forming a 'glial syncytium' (Mugnaini, 1986; Rash et al., 2001; Orthmann-Murphy et al., 2008). There are abundant gap junctions between astrocytes (A/A), fewer between oligodendrocytes and astrocytes (O/A), and few or none between oligodendrocytes themselves. Oligodendrocytes express Cx32 and Cx47 (Dermietzel et al., 1989; Micevych and Abelson, 1991; Scherer et al., 1995; Li et al., 1997; Menichella et al., 2003; Nagy et al., 2003a; Odermatt et al., 2003; Kamasawa et al., 2005), as well as Cx29, which does not appear to form gap junctions (Altevogt et al., 2002; Li et al., 2002; Altevogt and Paul, 2004; Kleopa et al., 2004); cf. (Nagy et al., 2003a). Astrocytes express Cx30 and Cx43 (Dermietzel et al., 1989; Yamamoto et al., 1990; Micevych and Abelson, 1991; Nagy et al., 1997; Kunzelmann et al., 1999; Nagy et al., 1999, 2001, 2003b; Rash et al., 2001). A/A coupling appears to be limited to homotypic channels [Cx43/Cx43 and Cx30/Cx30; (Swenson et al., 1989; Werner et al., 1989; Dahl et al., 1996)], but does not include Cx30/Cx43 heterotypic channels (Orthmann-Murphy et al., 2007b). O/A coupling is most likely mediated by Cx47/Cx43 and Cx32/Cx30 heterotypic channels (Orthmann-Murphy et al., 2007b).

In humans, Cx47/Cx43 channels appear to be essential for the proper maintenance of myelin. PMLD-associated mutations are recessive and result in the loss-of-function of Cx47, including the ability to form functional channels with Cx43, suggesting that the loss of O/A coupling mediated by Cx47/Cx43 channels causes PMLD (Orthmann-Murphy et al., 2007a, b). Here, we describe a novel mutation (I33M) in GJA12/GJC2 in three members of one family who have complicated hereditary spastic paraplegia (HSP). In a cell model system, the mutant protein forms gap junction plaques with Cx43, but I33M/Cx43 channels have such severely altered voltage dependent gating that they would not be predicted to function under physiological conditions.

Methods

MRI studies

MRI was performed with a 1.5 Tesla MR unit (Siemens Magnetom Avanto, Erlangen, Germany). The imaging protocol included sagittal T1-weighted spin-echo, axial T1-weighted inversion-recovery turbo spin-echo, axial proton-spin density and T2-weighted turbo spin-echo and coronal FLAIR. Single-section multivoxel 2D 1H MR spectroscopic imaging (1H MRSI) was acquired in all patients with a PRESS technique (repetition time (TR)/echo time (TE) = 1200/144 ms), with a nominal planar resolution of 1.04 mm x 24 x 24; field of view = 200 x 200 x 15 mm3). The volume of interest was positioned at the level of the centrum semiovale. Scan acquisition time was 14 min. 1H MRSI data were reconstructed and processed by using 'csx2' software (Soher et al., 1996). Imaging of the cervical spine with sagittal T1- and T2-weighted and axial T2-weighted MR images was acquired for patient III-8.

Mutation analysis

Oligonucleotide primers (Uhlenberg et al., 2004) were used to PCR-amplify the single coding exon of GJA12/GJC2 from genomic DNA in three overlapping fragments, which were sequenced as previously described (Bugiani et al., 2006).

Expression analysis of the I33M mutant

We generated the 99C>G (I33M) GJA12/GJC2 mutation from a human Cx47 cDNA sequence (GenBank accession number AF014643) using the QuikChange kit for PCR site-directed mutagenesis (Stratagene, La Jolla, CA, USA), with the following oligonucleotide primers (the underlined codon encodes the altered amino acid): 5’-gggaggctcttcgccatgtgtgcagcagctg-3’; 5’-cagcgctagcatacctggagaac/cacc-3’, as previously described (Orthmann-Murphy et al., 2007a). The resulting DNA [in pIRE2-EGFP and subcloned into pIRESpuro3 (Clontech, Mountain View, CA, USA)] and the mutation was confirmed by sequencing at the Sequencing Core of the University of Pennsylvania.
Neuro2A cells (N2A, from American Type Culture Collection, Manassas, VA, USA) and communication-incompetent HeLa cells (gift of Dr Klaus Willecke, University of Bonn, Bonn, Germany) were maintained and transfected to transiently or stably express mutant or WT Cx47 as previously described (Orthmann-Murphy et al., 2007a, b). Heterotypic mix experiments were performed on HeLa cells stably expressing WT Cx43, WT Cx47, I33M or P87S as described previously (Orthmann-Murphy et al., 2007b).

Cells were immunostained as previously described (Orthmann-Murphy et al., 2007a, b) using rabbit antibodies raised against the C-terminus of human Cx47 (diluted 1:1500; Orthmann-Murphy et al., 2007a) or mouse monoclonal antibodies against mouse Cx43 (diluted 1:1000; Millipore Corporation, Billerica, MA, USA) or pan-cadherin (Novus Biologicals, Littleton, CO, USA; diluted 1:200) and FITC-, TRITC-, and Cy5-conjugated secondary antibodies. Pan-cadherin (gift of Dr Klaus Willecke, University of Bonn, Bonn, Germany) was then imported into Adobe Photoshop (San Jose, CA, USA) then merged and pseudocoloured using Image J 1.37v (National Institutes of Health, USA) and communication-incompetent HeLa cells were collected and processed for immunoblotting as previously described (Orthmann-Murphy et al., 2007a).

Functional tests of the I33M mutant
Confluent HeLa cell monolayers were scrape loaded as previously described (Orthmann-Murphy et al., 2007a). We used dual whole-cell patch clamp to assess formation of both homotypic channels, on pairs of N2A cells transiently transfected with WT Cx43, WT Cx47 or I33M subcloned into pIRE2-EGFP, and heterotypic, on pairs of cells each expressing a pIRE2-EGFP construct or pIRE2-DsRed construct, as described previously (Orthmann-Murphy et al., 2007b). Recording solutions used were as follows (in mM): pipette solution, 145 CsCl, 5 EGTA, 0.5 CaCl₂ and 10.0 HEPES, pH 7.2; bath solution, 150 NaCl, 4 KCl, 1 MgCl₂, 2 CaCl₂, 5 dextrose, 2 pyruvate and 10 HEPES, pH 7.4. Heterotypic pairings between two cells are shown as ‘connexin expressed in cell 2/connexin expressed in cell 1’; junctional conductance (Gj)—junctional voltage (Vj) relations were determined from isolated pairs by measuring instantaneous junctional current (i) responses in cell 2 following 12.5-s Vj pulses (from −100 to 100 mV in 20 mV increments) applied to cell 1 and applying Ohm’s law. Baseline Gj was similarly determined by measuring instantaneous Ij responses to ±40 mV Vj pulses.

Statistical analyses
For heterotypic mix experiments, epifluorescence images were imported into Adobe Photoshop and analysed as previously described (Orthmann-Murphy et al., 2007b). At least 11 DsRed+ cells from two cover slips were analysed for each heterotypic mixture; this was repeated in three independent experiments. For each DsRed+ cell, we determined the mean and 95% CI of the number of overlapping cell surface puncta. The number of DsRed+ cells with or without one or more overlapping puncta was compared using Fisher’s exact test with Bonferroni’s correction for multiple comparisons. All statistical tests were performed in GraphPad Prism (San Diego, CA, USA). For dual whole-cell patch clamp assays, values are presented as mean Gj±SEM, and were compared using Fisher’s exact test or Mann–Whitney test in GraphPad Prism.

Results
Clinical findings
The family pedigree is shown in Fig. 1. There were three affected members—the proband (III-8, arrowhead), his brother (III-11) and their female cousin (III-5). At age 39, the proband presented with an 8-year history of slowly progressive walking difficulties, leg stiffness and slurred speech. He felt ‘awkward’ since his school years, had minimal writing difficulties, but was fit for military service at age 18. At age 21, he underwent surgery for concomitant strabismus. Currently, he is able to walk without assistance, and works as a priest. His legs and gait were spastic, and bilateral Babinski signs were found. The jaw jerk and deep tendon reflexes (DTRs) were brisk throughout, with non-sustained ankle clonus. Mild pes cavus was noted. He had no nystagmus, but pursuit eye movements were slightly saccadic. The following findings were mild: pseudobulbar dysarthria, loss of finger dexterity, dysmetria and intention tremor on finger-to-nose and heel-to-knee testing. The proband’s 36-year-old brother (III-11) had minimal motor difficulties since infancy and mild learning impairment at school. He reported mild intention tremor during late-childhood, and a few tonic-clonic seizures, mainly febrile, between childhood and his early teens. These problems had long been attributed to a dystocic delivery with cyanosis at birth. After age 20, he began to show a slow worsening of walking difficulties and slight dysarthria, followed by decreased dexterity with his hands. Presently, he walks with a cane, but he can still walk a few steps without aid. On neurological examination he had a shuffling, scissor-like gait, marked spasticity with mild weakness of the lower limbs, an increased jaw jerk and increased DTRs in all limbs, with sustained clonus at the ankles and bilateral Babinski signs. He had slight...
dysarthria, mild loss of finger dexterity, dysmetria, intention tremor on finger-to-nose testing and moderate hypopodiodochokinesia. He had pes cavus with hammer toes, lumbar hyperlordosis and concomitant strabismus but no nystagmus.

The proband’s 53-year-old cousin (III-5) had always been considered somewhat dull, but completed primary school without help. She walked at 20 months, and developed progressive spastic paraplegia and dysarthric speech beginning in her teens, becoming wheelchair bound at age 30. She later developed urinary incontinence followed by retention, and required a permanent catheter. Her jaw jerk and all DTRs were exaggerated, with sustained right ankle clonus and bilateral Babinski signs. She had moderate hand hypodiadochokinesia. Light touch and pain sensations were reduced in the left upper limb and both legs. Position and vibration sense were severely impaired in the legs. She also had severe dorsal scoliosis, bilateral pes cavus and ankle contractures (Table 1).

Clinical investigations

The following blood tests were normal for all affected patients: creatine kinase, uric acid, ammonia, lactate, pyruvate, alanine and other amino acids, very-long-chain fatty acids and phytic acid. Urinary levels of organic acids, amino acids and sulfatides were also normal. Lysosomal enzyme assessment in blood leukocytes ruled out metachromatic leukodystrophy, Krabbe disease and GM1 and GM2 gangliosidoses. Neuro-ophthalmologic examination revealed only left eye amblyopia in patients III-8 and III-11, attributed to congenital strabismus. Audiometry revealed mild sensorineural hearing loss in Patient III-5 and left ear conduction hypoacusia due to previous trauma in Patient III-8.

Electromyography and motor and sensory nerve conduction studies were normal in all three patients. Motor evoked potential (MEP) studies showed prolongation of central motor conduction time (CMCT) in upper limbs, and no response in lower limbs in Patients III-8 and III-11; in Patient III-5, MEPs could be recorded only from the left upper limb, with central motor conduction time prolongation, whereas no response was obtained from the other limbs. Their electroretinogram was normal, whereas visual evoked potentials showed delayed P100 wave latencies. Brainstem auditory evoked potentials disclosed absence or severe distortion and delay of waves III and V; wave I was normal in Patients III-11 and III-5, but almost unrecognizable in Patient III-8, particularly on the left side. Upper and lower limb somatosensory evoked potential studies demonstrated delayed latencies of central components in all patients; no response could be evoked in lower limbs in Patient III-5. In all patients, EEG showed a posterior background activity of about 8 Hz with a slightly irregular morphology. The cardiovascular reflexes were normal in Patient III-8. Neuropsychological assessment revealed IQ scores of 94 in Patient III-8, 83 in Patient III-11 and 77 in Patient III-5 (normal value > 70).

Table 1 Summary of clinical and instrumental data for patients with the I33M GJA12/GJC2 mutation

<table>
<thead>
<tr>
<th></th>
<th>Patient III-8</th>
<th>Patient III-11</th>
<th>Patient III-5</th>
</tr>
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<tbody>
<tr>
<td>Sex/Age at examination (years)</td>
<td>M/39</td>
<td>M/36</td>
<td>F/53</td>
</tr>
<tr>
<td>Age at disease onset</td>
<td>2nd decade</td>
<td>1st decade</td>
<td>1st decade</td>
</tr>
<tr>
<td>Age of disease progression (years)</td>
<td>&gt;30</td>
<td>&gt;20</td>
<td>&gt;10</td>
</tr>
<tr>
<td>Initial motor development</td>
<td>Normal</td>
<td>Normal</td>
<td>Walking at 2 yrs</td>
</tr>
<tr>
<td>Onset symptom</td>
<td>Difficulty in walking</td>
<td>Hand tremor</td>
<td>Difficulty in walking</td>
</tr>
<tr>
<td>Walking ability at exam</td>
<td>Without aid</td>
<td>With unilateral aid</td>
<td>Chairbound since age 30</td>
</tr>
<tr>
<td>Lower limb spasticity</td>
<td>++</td>
<td>+++</td>
<td>++++</td>
</tr>
<tr>
<td>Upper limb motor involvement</td>
<td>±</td>
<td>+</td>
<td>++</td>
</tr>
<tr>
<td>Dysarthria</td>
<td>+</td>
<td>++</td>
<td>++</td>
</tr>
<tr>
<td>Nystagmus</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Cerebellar ataxia</td>
<td>±</td>
<td>+</td>
<td>++</td>
</tr>
<tr>
<td>Mental impairment/QI score</td>
<td>±/94</td>
<td>±/83</td>
<td>+/77</td>
</tr>
<tr>
<td>Epilepsy</td>
<td>–</td>
<td>+</td>
<td>–</td>
</tr>
<tr>
<td>Sphincter dysfunction</td>
<td>–</td>
<td>–</td>
<td>++</td>
</tr>
<tr>
<td>Sensory loss</td>
<td>–</td>
<td>–</td>
<td>++</td>
</tr>
<tr>
<td>Sensorineural hearing loss</td>
<td>–</td>
<td>–</td>
<td>+</td>
</tr>
<tr>
<td>Other signs</td>
<td>Pes cavus, strabismus</td>
<td>Pes cavus, strabismus</td>
<td>Pes cavus, scoliosis</td>
</tr>
<tr>
<td>MRI</td>
<td>Hypomyelination</td>
<td>Hypomyelination</td>
<td>Hypomyelination and atrophy</td>
</tr>
<tr>
<td>1H-MRSI</td>
<td>↓ Cho/NAA, Cho/Cr ratios</td>
<td>↓ Cho/NAA, Cho/Cr ratios</td>
<td>↓ Cho/NAA, Cho/Cr ratios</td>
</tr>
<tr>
<td>EEG</td>
<td>Near-normal</td>
<td>Near-normal</td>
<td>Near-normal</td>
</tr>
<tr>
<td>EMG and nerve conduction studies</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
</tr>
<tr>
<td>VEPs, BAEPs, MEPs and SEPs</td>
<td>Abnormal</td>
<td>Abnormal</td>
<td>Abnormal</td>
</tr>
<tr>
<td>Autonomic tests</td>
<td>Normal</td>
<td>ND</td>
<td>ND</td>
</tr>
</tbody>
</table>

–= absent; ±= minimal; + = mild; ++ = moderate; +++ = marked; ++++ = severe; ND = not done. BAEPs = brainstem auditory evoked potentials; EMG = electromyography; MEPs = motor evoked potentials; SEPs = somatosensory evoked potentials; VEPs = visual evoked potentials.
Brain MR imaging and spectroscopic findings were homogeneous in the three patients and highly suggestive of a hypomyelinating leukoencephalopathy. White matter regions showed diffuse high signal intensity on T2-weighted and low signal on T1-weighted images (Fig. 2, Supplementary Fig. 1). T2-weighted signal hyperintensity was evident at level of the pons, in the region of corticospinal and spinothalamic tracts (Fig. 2B and F). The corpus callosum was thin in all patients. The cervical spine of Patient III-8 appeared normal (data not shown). Ventricular and posterior fossa dilation, likely secondary to white matter volume loss, was mild in Patients III-8 and III-11, but advanced in Patient III-5. Choline (Cho), N-acetyl-aspartate (NAA) and creatine (Cr) were measured in the white matter of the centrum semiovale by 1H MR spectroscopic imaging. The NAA/Cr ratios were within normal values, consistent with what has been reported for PMD and PMLD (Lee et al., 2004; Bizzi et al., 2008). Average Cho/NAA and Cho/Cr ratios, however, were reduced—0.49 and 0.84 in Patient III-8; 0.47 and 0.80 in Patient III-11; 0.44 and 0.75 in Patient III-5—compared with 0.60 and 1.0, respectively, in normal adults.

**Affected patients have an I33M mutation in GJA12/GJC2**

The above data indicated that the three patients from this family had a recessively inherited hypomyelinating leukoencephalopathy with an unusual clinical phenotype of a late onset complicated HSP. Although the parents of our patients were reported to be unrelated, they all originated from a small village in northern Italy, and Subjects I-2, I-3 and II-1 (Fig. 1) had the same family name. Hence, we sequenced amplified genomic DNA of the GJA12/GJC2 gene, and found a novel missense mutation (99C>G), predicted to cause an Ile>Met amino acid substitution (I33M) in Cx47. This mutation was homozygous in the three affected individuals, heterozygous in obligate healthy carriers and some clinically normal relatives (asterisks in Fig. 1 denote DNA analysis), and absent in 210 control alleles. Heterozygous relatives (Fig. 1) were asymptomatic and had a normal neurological exam (II-9, III-7 and III-12); one of these relatives (III-7) had a normal brain MRI and 1H MRSI.

As shown in Fig. 3, I33 is located in the first transmembrane domain, which is a highly conserved region of connexins (Yeager and Nicholson, 1996). This amino acid residue is identical in Cx47 orthologues of other vertebrates (cow, mouse, frog and zebrafish; Supplementary Fig. 2), but Val (Cx30, Cx30.3, Cx31, Cx31.1), Leu (Cx30.2) or even Met (Cx40, Cx59, Cx62) are found at the corresponding position in other connexins (Supplementary Fig. 2). Although mutations in the genes encoding Cx26, Cx30, Cx30.3, Cx31, Cx32, Cx40, Cx43, Cx46 and Cx50 have been described, mutations in the residues corresponding to I33 have not been identified previously.

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**Fig. 2** Imaging studies show white matter abnormalities. These are MRI images from patients III-8 (A–D) and III-5 (E–H). Sagittal T1-weighted (T1W) images [spin-echo (SE): repetition time (TR)/echo time (TE) 556/13 ms; 6-mm thickness] shows thinning of the corpus callosum, posteriorly (A, arrowhead) or diffusely (E). Axial T2W images (SE: TR/TE = 3200/90 ms; 5-mm thickness) show symmetric hyperintensity in the region of the corticospinal/corticobulbar tracts at the level of the pons B and F, arrowhead) and the posterior limb of internal capsule (C and G, arrowhead). In addition, there is diffuse hyperintensity in the subcortical, lobar and periventricular white matter (C, G and H), and enlarged ventricles, especially in Patient III-5 (G). Axial T1W image [inversion recovery turbo spin-echo (IRTE): TR/TE = 5600/70 ms; flip angle 150°; 4-mm thickness] shows diffuse hypointensity in the white matter, including the posterior limb of the internal capsule (D, arrowhead).
I33M forms gap junction plaques in HeLa cells

To investigate the underlying molecular defects of the I33M mutant, we expressed it in communication-incompetent HeLa and N2A cells. For comparison, we also expressed WT Cx47 and P87S, a missense mutant that causes PMLD (Uhlenberg et al., 2004), and results in loss-of-function (Orthmann-Murphy et al., 2004), and in bulk-selected and transiently transfected cells (Supplementary Fig. 3). As expected (Orthmann-Murphy et al., 2007a, b), P87S was localized to the endoplasmic reticulum (ER) in both transiently (Supplementary Figs. 4 and 5) and permanently transfected (Fig. 4) cells, whereas I33M (and WT Cx47) formed gap junction plaques at apposed cell borders. We confirmed the cell surface localization of WT Cx47 and I33M by double labelling with a monoclonal antibody against pan-cadherin (green) and counterstained with DAPI. The pan-cadherin staining at cell borders interdigitates with the cell surface staining of Cx47 in cells that express WT Cx47 (arrowheads) or I33M (arrowheads), but surrounds the staining of cells expressing the mutant P87S, which is localized in the endoplasmic reticulum. Scale bar: 10 μm.

I33M does not form functional homotypic channels

Because patients expressing I33M have a milder phenotype than do patients with PMLD, and I33M can form gap junction plaques,
we hypothesized that I33M might form gap junction channels with altered functional properties. To determine whether I33M could transfer small molecules, we scrape loaded cells (el-Fouly et al., 1987; Trosko et al., 2000), using gap junction tracers of different sizes, shapes and charge. A confluent monolayer of cells was injured with a scalpel blade in media that contained 2% neurobiotin (NB; MW 287, +1) or 0.1% Lucifer Yellow (LY; MW 443, -2). As in parental cells or cells expressing vector alone (data not shown), no transfer was seen past the scrape line in bulk-selected cells expressing I33M or P87S (Fig. 5B–C and E–F), whereas NB and LY transferred to cells beyond the scrape line for cells expressing WT Cx47 (Fig. 5A and D). We also confirmed that the cells along the scrape line expressed the appropriate connexin (Supplementary Fig. 6). This experiment was repeated at least twice in two different bulk-selected cells lines expressing I33M with similar results.

We used a more sensitive assay to determine whether Cx47 mutants can form functional homotypic gap junctions—dual whole-cell patch clamping on transiently transfected N2A cells. In this experiment, each cell was transiently transfected to express a single connexin as well as EGFP. For negative control pairs, cells expressing a single connexin were paired with cells expressing EGFP alone (Cx/EGFP). Voltage ramps or steps were applied to one cell of an EGFP-expressing pair, and current responses were measured in the second cell; the junctional voltage (V) corresponds to the difference in voltage between the two cells. As shown in Table 2, I33M/I33M homotypic pairings failed to form functional channels (0/6 pairs), whereas cell pairs expressing

Fig. 5 I33M is impermeable to low molecular weight tracers. These are representative images of bulk-selected HeLa cells that stably express WT Cx47 or the indicated mutants, scrape-loaded with neurobiotin (A–C) or Lucifer Yellow (D–F). Cells scrape-loaded with neurobiotin were fixed with 4% paraformaldehyde, then visualized using FITC-conjugated extravidin (green) and DAPI counterstain (blue). Cells scrape-loaded with Lucifer Yellow were imaged by epifluorescence (top row) and phase contrast (bottom row) 5 min after scrape loading. Only HeLa cells expressing WT Cx47 showed transfer of neurobiotin (A) or Lucifer Yellow (D) to neighbouring cells. The images for each type of scrape loading experiment were acquired on the same day at the same exposure. (A–C) scale bar: 20 μm; (D–F) scale bar: 20 μm.
WT Cx47 were coupled. The difference between I33M/I33M and WT Cx47/Cx47 pairings was statistically significant (I33M/I33M versus Cx47/Cx47, P=0.0022, Fisher’s exact test). Thus, I33M does not appear to form functional homotypic gap junctions by two different assays.

**Table 2** Summary of dual whole-cell patch clamp recordings

<table>
<thead>
<tr>
<th>n</th>
<th>Mean Gj (nS)</th>
<th>SD</th>
<th>SEM</th>
</tr>
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<tbody>
<tr>
<td>I33M/I33M</td>
<td>6</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>I33M/Cx47</td>
<td>15</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Cx47/Cx47</td>
<td>5</td>
<td>19.31</td>
<td>23.58</td>
</tr>
<tr>
<td>I33M/Cx43</td>
<td>13</td>
<td>0.05</td>
<td>0.13</td>
</tr>
<tr>
<td>Cx47/Cx43</td>
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<td>29.12</td>
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<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Cx47/EGFP</td>
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</tr>
<tr>
<td>Cx43/EGFP</td>
<td>3</td>
<td>0.50</td>
<td>0.87</td>
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</tbody>
</table>

N2A cells transiently expressing WT Cx43, WT Cx47 or I33M were paired as noted; using dual whole-cell patch clamping, junctional conductance (Gj) was measured when junctional voltage (Vj) was zero. No coupling was detected between pairs of cells either each expressing I33M (I33M/I33M) or separately expressing I33M or WT Cx47 (I33M/Cx47). Extremely low levels of coupling were detected between pairs of cells separately expressing I33M or WT Cx43 (I33M/Cx43). SEM=standard error of the mean; DsRed=pIRES2-DsRed monomeric to the ER, and did not appear to form overlapping puncta.

I33M forms functional heterotypic channels with Cx43 in a cell model system

To determine whether I33M can form functional channels with Cx43, we applied morphological and functional assays we used previously to show that Cx47 mutants associated with PMLD do not form functional heterotypic gap junctions with WT Cx43 (Orthmann-Murphy et al., 2007). In the morphological assay, cell lines in which at least 90% of cells stably expressed Cx47 (Orthmann-Murphy et al., 2007b). In contrast, we detected Gj activation for six of 13 (46%) I33M/Cx43 pairs tested, somewhat less than we previously found for all of the WT Cx47/Cx43 heterotypic pairs we have tested (40 of 55; 73%). As shown in Table 2, the mean Gj measured at Vj = 0 for I33M/Cx43 channels is significantly smaller than that of WT Cx47/Cx43 channels (I33M/Cx43 versus WT Cx47/Cx43, P<0.05, Mann–Whitney test). Negative control pairs, in which cells expressing a single connexin were paired with cells expressing EGFP or DsRed alone, showed low levels of coupling probably due to formation of heterotypic channels with an endogenous connexin expressed by parental N2A cells (Table 2; Orthmann-Murphy et al., 2007b).

To better define the functional differences between I33M/Cx43 and Cx47/Cx43 channels, we examined macroscopic current responses and Gj–Vj relations. By convention, pairing designation is ‘connexin expressed by cell 2/connexin expressed by cell 1’. Both cells in the pair were voltage clamped to 0 mV; cell 1 was stepped between −100 and 100 mV in 20 mV increments, and current was recorded from cell 2. For I33M/Cx43 heterotypic pairings, negative pulses (−40 mV) applied to the cell expressing Cx47 activate junctional currents (Ij) (Fig. 7A). Assuming that Gj is a linear function of Vj between −20 mV and +20 mV, the normalized Gj versus junctional voltage (Vj) relation reveals that I33M/Cx43 channels are about fifty times more likely to be open after a 12.5-s pulse to Vj = −100 mV than at Vj = 0 mV (solid line, Fig. 7B). This is strikingly different than WT Cx47/Cx43 channels (dashed line), which are more likely to be open at Vj = 0 mV than at either −100 mV or +100 mV (Fig. 7B; Orthmann-Murphy et al., 2007b).

The left shift of the Gj–Vj relation for I33M/Cx43 channels is likely explained by an extremely low open probability of the I33M hemichannel when Vj<−40 mV are applied to the cell expressing Cx43; this can account for the significantly reduced Gj values detected at Vj=0 mV for both I33M/Cx43 and I33M/Cx43 channels (Table 2). Although the WT Cx43 hemichannel has negative Vj gating polarity (Bukauskas et al., 2001) and should be closed under the conditions that open I33M hemichannels, Cx43 rarely closes fully (Bukauskas et al., 2000). It is possible, then, that I33M is forming a channel with a substate of Cx43. Properties of I33M/WT Cx47 channels (5/15 pairs) are similar to those described for I33M/Cx43 (data not shown).
Discussion

The I33M Cx47 mutant causes hereditary spastic paraplegia

The three patients with a homozygous 99C>G mutation in GJA12/GJC2 all developed HSP, characterized by a progressive gait disorder during the first to fourth decades of life. The proband exhibited an almost pure spastic paraplegia with mild ataxia and dysarthria, and walked unaided at age 39. His brother had moderately complicated spastic paraplegia, with mild ataxia and dysarthria, a few seizures and walked with unilateral support at age 36. Their cousin had a more severe and complex phenotype, with earlier onset and became wheelchair dependent at age 30. The late onset, preserved ability to walk unaided until adulthood, and
absence of nystagmus clearly distinguish these patients from the more severe PMLD phenotype caused by other recessive mutations in GJA12/GJC2 (Fig. 3). These PMLD patients typically present by 12 months of age with nystagmus and impaired psychomotor development, and rarely walk unaided in childhood (Uhlenberg et al., 2004; Bugiani et al., 2006; Salviati et al., 2007; Wolf et al., 2007; Henneke et al., 2008).

Like PLP1, mutations in GJA12/GJC2 appear to cause a spectrum of CNS white matter disease, including HSP and PMD/PMLD. Based on the similar phenotypes caused by PLP1 and GJA12/GJC2 mutations, one suspects that other causes of the PMLD (and HSP with primarily white matter pathology) will turn out to have altered 1H MRSI metabolic profiles in the three patients homozygous for the GJA12 mutation suggested a hypomyelinating leukoencephalopathy, which is consistent with oligodendrocyte expression of Cx47. In our adults patients, the T2-signal abnormalities were more subtle than those shown for previously reported PMLD patients (caused by GJA12/GJC2 mutations), who ranged in age from 5 months to 12 years (Uhlenberg et al., 2004; Bugiani et al., 2006; Salviati et al., 2007; Wolf et al., 2007). Notably, T2-signal hyperintensity for patients with the GJA12 mutation was discrete and prominent in the corticospinal and spinothalamic tracts (especially at the level of the pons). A similar involvement of these tracts is evident in the MR images of spastic PMLD patients reported by Wolf and colleagues (2007), whereas the corticospinal tracts appeared to be relatively spared in PMLD patients with other GJA12/GJC2 mutations (Bugiani et al., 2006). In our patients, the involvement of the long white matter tracts may correlate with the severity of spastic paraplegia. The mild decrease in metabolite ratios (Cho/NAA and Cho/Cr) measured in our patients, and previously reported for one of two PMLD patients with GJA12/GJC2 mutations (Bugiani et al., 2006), are in agreement with a recent study suggesting that hypomyelinating disorders (including PMLD due to GJA12/GJC2 mutations) are associated with normal or near-normal metabolite ratios, compared to a large increase in choline or decrease in NAA typically seen in actively demyelinating leukoencephalopathies (Bizzi et al., 2008). Finally, like PMLD patients with GJA12/GJC2 mutations, our patients exhibited central conduction slowing without electrophysiological evidence for peripheral neuropathy.

Hereditary spastic paraplegias are a heterogeneous group of genetic disorders characterized by progressive lower limb spasticity (pure HSP) that may be associated with other neurological abnormalities (complicated HSP; Fink, 2006). We propose that autosomal recessive mutations in GJA12/GJC2 that cause HSP should be termed SPG44 (http://www.genenames.org). The patients described here have a slowly progressive disease consisting of a spastic paraplegia; the inclusion of minimal to moderate ataxia and dystarthis, and seizures suggest that they have a complicated HSP. Although some of the clinical findings indicate that the pathophysiology is not limited to the longest CNS axons, other forms of ‘complicated’ HSP have comparable, or even more severe, findings (http://www.ncbi.nlm.nih.gov/sites/entrez?db=OMIM). Similarly, MRI abnormalities in the white matter, comparable to the ones we report, have been found in one patient with SPG2 (Lee et al., 2004). Thus, GJA12/GJC2 is only the second example (PLP1 is the other) of mutations that cause HSP in a gene that is primarily expressed by oligodendrocytes; the other HSP-causing loci are thought to primarily affect neurons/axons (Fink, 2006). Screening for GJA12/GJC2 mutations, therefore, should be considered for patients and/or families that present with a complicated HSP phenotype and hypomyelinating leukoencephalopathic findings on MRI.

The diffuse white matter MR signal abnormalities and mildly altered 1H MRSI metabolic profiles in the three patients homozygous for the GJA12 mutation suggested a hypomyelinating leukoencephalopathy, which is consistent with oligodendrocyte expression of Cx47. In our adults patients, the T2-signal abnormalities were more subtle than those shown for previously reported PMLD patients (caused by GJA12/GJC2 mutations), who ranged in age from 5 months to 12 years (Uhlenberg et al., 2004; Bugiani et al., 2006; Salviati et al., 2007; Wolf et al., 2007). Notably, T2-signal hyperintensity for patients with the GJA12 mutation was discrete and prominent in the corticospinal and spinothalamic tracts (especially at the level of the pons). A similar involvement of these tracts is evident in the MR images of spastic PMLD patients reported by Wolf and colleagues (2007), whereas the corticospinal tracts appeared to be relatively spared in PMLD patients with other GJA12/GJC2 mutations (Bugiani et al., 2006). In our patients, the involvement of the long white matter tracts may correlate with the severity of spastic paraplegia. The mild decrease in metabolite ratios (Cho/NAA and Cho/Cr) measured in our patients, and previously reported for one of two PMLD patients with GJA12/GJC2 mutations (Bugiani et al., 2006), are in agreement with a recent study suggesting that hypomyelinating disorders (including PMLD due to GJA12/GJC2 mutations) are associated with normal or near-normal metabolite ratios, compared to a large increase in choline or decrease in NAA typically seen in actively demyelinating leukoencephalopathies (Bizzi et al., 2008). Finally, like PMLD patients with GJA12/GJC2 mutations, our patients exhibited central conduction slowing without electrophysiological evidence for peripheral neuropathy.

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Like PLP1, mutations in GJA12/GJC2 appear to cause a spectrum of CNS white matter disease, including HSP and PMD/PMLD. Based on the similar phenotypes caused by PLP1 and GJA12/GJC2 mutations, one suspects that other causes of the PMLD (and HSP with primarily white matter pathology) will turn
out to be caused by mutations in genes that primarily affect oligodendrocytes. One example of this may be mutations of HSPD1 (the gene encoding the mitochondrial chaperonin, Hsp60), which cause either uncomplicated autosomal dominant SPG13 (Hansen et al., 2002) or an autosomal recessive lethal disease similar to conntnnal PMD (Magen et al., 2008).

The molecular pathogenesis of HSP and PMLD due to mutations in GJA12/GJC2

Because the patients with homozygous I33M mutations have a milder phenotype than do patients with PMLD, we predicted that I33M would have a partial loss-of-function, as compared to the failure to detect I33M/I33M dye coupling or currents and the formation gaps between astrocytes, we predicted that I33M would still form functional, but likely altered, channels with Cx43. The finding that I33M formed gap junction plaques with itself and with Cx43 is consistent with this hypothesis, and demonstrates that I33M oligomersizes into hemichannels that traffic to the cell surface (Kumar and Gilula, 1996), compared with the ER-retained Cx47 mutants which cause PMLD. The failure of I33M to transfer small molecules such as Lucifer Yellow or neurobiotin (in scrape loading assays), or to form detectable homotypic channels (by dual whole-cell patch clamping), argues against this hypothesis. Both the failure to detect I33M/I33M dye coupling or currents and the alterations in the I33M/Cx43 $G_{j}-V_{j}$ relation are likely due to a shift of the open probability of the I33M hemichannel as a function of voltage, such that its open probability increases only when a large voltage difference ($V_{j} \leq -40$ mV) is applied with the respect to the cell expressing Cx43. Some disease-associated Cx32 mutants show similar alterations in voltage dependence and are also thought to lead to loss-of-function (Oh et al., 1997; Abrams et al., 2001). Because large voltage gradients that open I33M/Cx43 channels probably do not occur across O/A junctions, the I33M mutant should cause loss-of-function of these channels. Large O/A gradients would be particularly improbable if these cells are well coupled by Cx30/Cx32 channels (Orthmann-Murphy et al., 2007b, 2008). Consequently, similar to patients with GJA12/GJC2 mutations that cause PMLD, it is unlikely that Cx47/Cx43 gap junctions contribute to O/A coupling in patients with homozygous I33M mutations.

If the I33M mutation completely disrupts gap junction coupling via Cx47/Cx43 channels like the Cx47 mutants that cause PMLD, then another mechanism must account for the milder phenotype in our three patients. One possibility is that Cx47 has functions in oligodendrocytes beyond forming functional O/A channels. Because Cx47 has a PDZ binding domain [and at least one binding partner, ZO-1; (Li et al., 2004)], it is possible that I33M, in particular, retains some important function (such as binding ZO-1) that requires proper localization at the cell surface. Along these lines, it is possible that like some other connexins (Prochnow and Dermietzel, 2008), Cx47 is required for cell adhesion, as shown for Cx26 and Cx43 in neuronal migration in developing cortex (Elias et al., 2007), and Cx32 and Cx43 in an in vitro aggregation assay (Cotrina et al., 2008). Another possibility is that the recessive Cx47 mutants that cause PMLD (P87S, Y269D, M283T) actually have dominant effects (that the I33M mutant does not have) that contribute to the more severe phenotype of PMLD. Because the P87S, Y269D and M283T mutants are mostly found in the ER, it seems unlikely that they would form an abnormal hemichannel on the cell membrane (Richardson et al., 2004; Dobrowolski et al., 2007). Alternatively, because they accumulate in the ER, these mutants could induce an unfolded protein response with deleterious effects, but we found no evidence for this possibility in a cell model system (Orthmann-Murphy et al., 2007a).

Finally, it is possible that Cx47 mutants associated with more severe phenotypes result in more axonal degeneration, independent of their effects on myelin. In the peripheral nervous system, mutations in genes expressed by Schwann cells (such as GJB1, PMP22 and MPZ) cause demyelinating neuropathy; although the demyelination is Schwann cell autonomous, clinical disability correlates with the degree of axonal damage (Nave et al., 2007; Scherer and Wrabetz, 2008). The enlarged ventricles in Patient III-5 (who was the most severely affected) are likely a result of axonal loss; such secondary axonal degeneration probably contributes to progressive neurological deterioration in PMD (Inoue, 2005; Garbern, 2007), and multiple sclerosis (Bjartmar et al., 1999). In spite of its importance, it is unclear how axons are damaged in any of these demyelinating disorders; in the case of Cx47 mutants, it is possible that disrupted O/A coupling impedes K$^{+}$ buffering mediated by glial gap junctions (Menichella et al., 2006).

Supplementary material

Supplementary material is available at Brain online.

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References


Altevogt BM, Kleopa KA, Postma FR, Scherer SS, Paul DL. Cx29 is uniquely distributed within myelinating glial cells of the central and peripheral nervous systems. J Neurosci 2002; 22: 6458–70.


GJA12 mutation causes spastic paraplegia


Nagy JJ, Ionescu AV, Lynn BD, Rash JE. Coupling of astrocyte connexins Cx26, Cx30, Cx43 to oligodendrocyte Cx29, Cx32, Cx47: implications from normal and connexin32 knockout mice. Glia 2003b; 44: 205–18.


