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Post-translational modifications of connexin26 revealed by mass spectrometry

Darren LOCKE*1,3 , Shengjie BIAN*1,2 , Hong LI† and Andrew L. HARRIS*

*Department of Pharmacology and Physiology, New Jersey Medical School, University of Medicine and Dentistry of New Jersey, Newark, NJ 07103, U.S.A., and †Center for Advanced Proteomics Research, New Jersey Medical School Cancer Center, University of Medicine and Dentistry of New Jersey, Newark, NJ 07103, U.S.A.

Gap junctions play important roles in auditory function and skin biology; mutations in the Cx26 (connexin26) gene are the predominant cause of inherited non-syndromic deafness and cause disfiguring skin disorders. Mass spectrometry (MS) was used to identify PTMs (post-translational modifications) of Cx26 and to determine whether they occur at sites of disease-causing mutations. Cx26 was isolated from transfected HeLa cells by sequential immunoaffinity and metal chelate chromatography using a tandem C-terminal haemagglutinin epitope and a (His-Asn)₆ sequence. In-gel and in-solution enzymatic digestions were carried out in parallel with trypsin, chymotrypsin and endoproteinase GluC. Peptides were fractionated using a reversed-phase matrix by stepwise elution with increasing concentrations of organic solvent. To improve detection of low-abundance peptides and to maximize sequence coverage, MALDI-TOF-MS (matrix-assisted laser desorption/ionization-time-of-flight mass spectrometry; MS) and MALDI-TOF/TOF-MS/MS (matrix-assisted laser desorption/ionization-time-of-flight/time-of-flight tandem mass spectrometry; MS/MS) spectra were acquired from each elution

INTRODUCTION

At least four connexin isoforms are expressed in the cochlea. Prominent among those, Cx26 (connexin26) is found in the supporting cells of the organ of Corti, the basal cell region of the stria vascularis and type I fibrocytes of the spiral ligament, which are important components for auditory function [1]. Mutations in the Cx26 gene (GJB2) are the predominant cause of inherited non-syndromic sensorineural deafness in humans [2,3]. Knockout mouse studies have provided insights into the role of Cx26 in auditory function and have demonstrated that Cx26 is essential for cochlear function and survival of the sensory epithelium of the inner ear [4,5]. Although Cx26 mutations that cause complete loss of functional junctional channels lead to non-syndromic deafness, which is principally recessive, mutations that cause altered channel function can lead to dominant deafness syndromes that include severe and disfiguring skin pathologies [6]. These mutations are found in all topological domains of Cx26; connexin proteins have four transmembrane domains (M1-M4), cytoplasmic N-terminal (NT) and C-terminal (CT) domains, one cytoplasmic loop (CL) between M2 and M3, and two extracellular loops (E1, between M1 and M2, and E2, between M3 and M4).

step using an Applied Biosystems 4800 tandem mass spectrometer. Acquisition, processing and interpretation parameters were optimized to improve ionization and fragmentation of hydrophobic peptides. MS and MS/MS coverage of Cx26 was significantly above that reported for other membrane proteins: 71.3 % by MS, with 29.9 % by MS/MS. MS coverage was 92.6 % if peptides resulting from in-source collisions and/or partial enzymatic cleavages were considered. A variety of putative PTMs of Cx26 were identified, including acetylation, hydroxylation, γ carboxyglutamation, methylation and phosphorylation, some of which are at sites of deafness-causing mutations. Knowledge of the PTMs of Cx26 will be instrumental in understanding how alterations in the cellular mechanisms of Cx26 channel biogenesis and function lead to losses in auditory function and disfiguring skin disorders.

Key words: connexin, deafness, mass spectrometry, mutation, post-translational modification, skin disease.

 $Cx26_{deaf}$ (mutations of Cx26 that cause non-syndromic or syndromic human deafness) disrupt the intercellular molecular/ionic signalling pathway by affecting connexin channel function, assembly and/or trafficking [7]. In other proteins, these processes can be modulated or controlled by PTMs (post-translational modifications) [8]. With the exception of phosphorylation, predominantly studied in Cx43 (connexin43), modulation of connexin channel properties and cellular dynamics by PTMs have not been studied. Furthermore, the potential role of disrupted PTMs in Cx26_{deaf} pathologies is unexplored. Knowledge of Cx26 PTMs will be instrumental in guiding experiments to understand how cellular mechanisms of connexin channel biogenesis and function can become altered, and how Cx26_{deaf} mutation can lead to losses in auditory function and to skin pathology.

Unfortunately, analysis of PTMs of membrane proteins by mass spectrometry (MS) remains challenging. Sequence coverage of membrane proteins is typically lower than that of soluble proteins, largely because of proteolytic resistance, and poor ionization of hydrophobic peptides makes detection of low-abundance peptides in complex spectra and PTM identification by tandem MS (MS/MS) difficult [9,10].

Abbreviations used: a.m.u., atomic mass units; CAM-Cys, carboxyamidomethylcysteine; Chymo, chymotrypsin; CI, confidence interval; CID, collisioninduced dissociation; CL, cytoplasmic loop; CT, C-terminal domain; Cx26, connexin26; Cx26_{deaf}, mutations of Cx26 that cause non-syndromic or syndromic human deafness; DoDM, *n*-dodecyl-β-n-maltoside; E1 (etc.), extracellular loop 1 domain (etc.); ESI-LC-QTOF, electrospray ionization liquid chromatography coupled with quadrupole time-of-flight; GluC, endoproteinase GluC; HA, haemagglutinin; LC, liquid chromatography; M1 (etc.), transmembrane 1 domain (etc.); MALDI, matrix-assisted laser desorption/ionization; MS, mass spectrometry; MS/MS, tandem MS; NT, N-terminal domain; OG, *n*-octylβ-n-glucopyranoside; Oxi-Met, methionine oxidation; PTM, post-translational modification; QTOF, quadrupole time-of-flight; SCX, strong cation exchange; TCEP, tris-(2-carboxyethyl)phosphine; TFA, trifluoroacetic acid; TOF, time-of-flight; Tryp, trypsin.

¹ These authors contributed equally.

² Present address: Department of Chemical Biology, Rutgers University, Piscataway, NJ 08854, U.S.A.

³ To whom correspondence should be addressed (email lockeda@umdnj.edu).

In the present study, significant methodological improvements were made to existing methods for MALDI–TOF-MS (matrix-assisted laser desorption/ionization–time-of-flight mass spectrometry) and were used to identify Cx26 PTMs. Cx26 was isolated by sequential immunoaffinity and metal chelate chromatography using a pair of tandem C-terminal tags. In-gel and in-solution digestions were carried out in parallel with three enzymes, with peptides recovered with and fractionated from a reversed-phase matrix to improve the recovery of low-abundance peptides in complex digestion mixtures, and thereby maximize sequence coverage. MS acquisition, processing and interpretation parameters were also optimized to improve ionization and MS/MS fragmentation of hydrophobic connexin peptides.

The results reveal a variety of PTMs of Cx26, some of which are at $Cx26_{deaf}$ sites. The findings are interpreted in the context of what is known about the effects of mutations at those sites on connexin channel biogenesis and function.

EXPERIMENTAL

Materials

Components of the Tet-On connexin expression system were from BD Biosciences. DMEM (Dulbecco's modified Eagle's medium), G418 sulfate, hygromycin and doxycycline were from Life Technologies. Agarose-conjugated anti-HA (haemagglutinin) clone HA-7 mouse IgG was from Sigma, as were other reagents unless stated. Talon superflow metal (cobalt) affinity resin was from Clontech. The detergent OG (*n*-octyl- β -D-glucopyranoside; 99.5 % purity) was from Glycon Biochemicals, and DoDM (*n*-dodecyl- β -D-maltoside; \geq 95 % purity) was from EMD Biosciences. Water was LC (liquid chromatography)-MS grade or quartz-distilled reverse-osmosis-purified.

Expression and purification of Cx26 channels

Bidirectional tetracycline-responsive expression vectors (Clontech) were used to express Cx26 channels in HeLa cells, which have virtually no endogenous connexin expression [11]. The rat Cx26 coding sequence was subcloned in-frame with a sequence coding for a 3.3 kDa CT domain 'tag', consisting of a thrombin cleavage site followed by an HA epitope and six His-Asn repeats. Tet-On cells were maintained in 200 μ g/ml hygromycin and 100 μ g/ml G418. Cells were induced for Cx26 expression when approx. 35% confluent with 1 μ g/ml doxycycline for 48 h [12].

Tagged homomeric Cx26 hemichannels were purified from confluent Tet-On HeLa cells (8000-10000 cm²), solubilized in 50 mM NaH₂PO₄, 50 mM NaCl, 5 mM EDTA, 5 mM EGTA, 1 % (w/v) DoDM, 1 mM 2-mercaptoethanol, 0.5 mg/ml azolectin and 0.5 mM iPr₂*P*-F (di-isopropyl fluorophosphate; Calbiochem), pH 7.4, with added protease and phosphatase inhibitors (cocktails I and II; Sigma), for 2 h at 4°C with rocking. The supernatant $(100\,000\,\mathbf{g}_{av}$ for 30 min at 4 °C) was incubated with 0.25 ml of agarose-immobilized anti-HA mouse IgG overnight at 4°C with shaking. The antibody matrix was collected (700 g_{av} for 1 min at 4°C), and washed in a fritted column with 10 ml of 0.1 M PBS, 1 M NaCl and 0.5 % DoDM, pH 7.4, followed by 15 ml of the same buffer containing 138 mM NaCl. Bound Cx26 was eluted with 50 mM sodium acetate, 0.5 M NaCl, 10 mM KCl and 0.2 % DoDM, pH 4.0, and 0.6 ml fractions were collected into 0.05 ml of 1 M NaHCO₃, 10 mM KCl and 0.2 % DoDM, pH 9.0. The final pH was approx. 7.4. Solubilization of gap junctions with the glucoside detergents DoDM and OG yields hemichannels [13].

Elution fractions containing Cx26 (as determined by Western blot analysis with a monoclonal anti-HA antibody) were combined and diluted 5-fold with 0.1 M PBS, pH 7.4, containing 0.2 % DoDM and EDTA-free protease and phosphatase inhibitors (Roche). Cx26 was repurified by immobilized metal affinity chromatography using 0.5 ml of Talon superflow cobalt affinity resin. After incubation for 2 h at 4 °C, the resin was washed in a fritted column with 5 ml of 0.1 M PBS, pH 7.4, 10 mM imidazole and 0.2 % DoDM, then Cx26 was eluted by 0.1 M PBS, pH 7.4, 0.1 M imidazole and 0.2 % DoDM. After 0.1 % TFA (trifluoroacetic acid) was added to the eluate, purified Cx26 was concentrated by centrifugation (3800 g_{av} for 20 min at 25 °C) using an AmiconUltra4 30 kDa cut-off filter (Millipore).

SDS/PAGE, Western blot analysis and gold staining

Cx26 was resolved by SDS/PAGE on 12.5 % (w/v) acrylamide Tris/HCl gels (Bio-Rad Laboratories) under denaturing and reducing conditions. For Western blot analysis, proteins were transferred on to PVDF membranes and stained with specific antibodies to Cx26 (Zymed) or the HA epitope of the CT domain tag. Reactive species were detected with the relevant phosphatase-conjugated secondary antibodies using NBT/BCIP (Nitro Blue Tetrazolium/5-bromo-4-chloroindol-3-yl phosphate; Pierce Endogen). For gold staining, nitrocellulose membranes were used, blocked (0.1 M PBS, pH 7.4, and 0.3 % Tween 20; 30 min for 37 °C), washed and stained with colloidal gold total protein stain (Bio-Rad Laboratories). The blots were washed with quartz-distilled reverse-osmosis-purified water and air-dried.

In-gel digestion

Cx26, resolved by SDS/PAGE, was visualized with E-Zinc stain (Pierce Endogen). The Cx26 band was excised with a sterile scalpel blade and cut into 1 mm³ pieces, washed with 50% (v/v) acetonitrile, destained according to the manufacturer's instructions and dehydrated (10 min) in 100% acetonitrile. Gel pieces were rehydrated (10 min) in 0.1 M NH₄HCO₃, dehydrated (10 min) in 100% acetonitrile, and reduced with 0.1 M TCEP [tris-(2-carboxyethyl)phosphine] in 0.1 M NH₄HCO₃ (30 min at 37 °C). Following 'washing' by dehydration–rehydration–dehydration as above, alkylation was performed with 50 mM iodoacetamide in 0.1 M NH₄HCO₃ [1 h at room temperature (25 °C) in the dark]. Gel pieces were then 'washed' as described previously.

Sequencing-grade Tryp (trypsin; $1 \mu g/\mu l$ in 50 mM acetic acid; Stratagene), modified to resist autodigestion, and Chymo (chymotrypsin; $1 \mu g/\mu l$ in 50 mM acetic acid; Roche) were diluted 1/20 with 50 mM NH₄HCO₃. Gel pieces were rehydrated in a minimal volume of 50 mM NH₄HCO₃ and digested overnight at 37 °C with 50–100 ng of Tryp or Chymo. Sequencing-grade GluC (endoproteinase GluC; $1 \mu g/\mu l$ in water; Roche) was diluted 1/20 with 50 mM NH₄HCO₃. Gel pieces were rehydrated in a minimal volume of 50 mM NH₄HCO₃ and digested overnight at 25 °C with 50–100 ng of GluC.

After overnight digestion, Cx26 peptides from each digestion were extracted with $10-20 \ \mu$ l suspensions of 2.5 % (v/v) Poros-20R2 reversed-phase beads (Applied Biosystems) in 5 % (v/v) acetic acid, 0.2 % TFA and 0.1 % OG by vortex-mixing (1000 rev./min at 4 °C) overnight.

Poros-20R2 beads were back-loaded on to micro- C_{18} or micro-SCX (strong cation exchange) ziptips (Millipore) and were washed according to the manufacturer's instructions. Cx26 peptides were recovered by step gradient elution: for C_{18}

ziptips, elution conditions were 5, 15, 25, 35, 60 and 80% (v/v) acetonitrile, followed by 50% (v/v) methanol/20% (v/v) acetonitrile (all in 0.1% TFA); for SCX ziptips, elution conditions were 1, 5 and 10% (v/v) NH_4OH in 30% (v/v) methanol [13].

Solution volumes were reduced by brief drying under vacuum before each elution fraction was mixed with matrix/standard solution {7 mg/ml α -cyanohydroxycinnamic acid in 50% (v/v) methanol, 20% (v/v) acetonitrile and 0.1% TFA containing 50 fmol/ μ l each of proteomics grade Glu-fibrinogen peptide [*m*/*z* 1570.677 a.m.u. (atomic mass units)] and ACTH_{18–39} (*m*/*z* 2465 a.m.u.) (both from AnaSpec)}. Aliquots (0.5–1.0 μ l) were spotted on to a stainless steel MALDI plate and were analysed with a 4800 proteomics analyser tandem mass spectrometer (Applied Biosystems).

In-solution digestion

Purified concentrated Cx26 was reduced with 0.1 M TCEP (for 30 min at 37 °C in the dark), and then immediately alkylated with 50 mM iodoacetamide (1 h at room temperature in the dark), both in 0.1 M NH₄HCO₃. Cx26 samples (1 vol.) were precipitated by phase separation using 2 vols of ice-cold methanol, and 1 vol. of ice-cold chloroform, with vortex-mixing after each addition. For phase separation, 1.5 vols of ice-cold water were added, the samples were vortex-mixed and centrifuged (10000 g_{av} for 5 min at 4 °C). The upper phase was discarded and 1.5 vols of cold methanol was added to the lower phase and protein/Cx26 at the interface. The samples were vortex-mixed and centrifuged (14000 g_{av} for 5 min at 4 °C) to pellet protein/Cx26. Excess solution was removed and the pellet was dried briefly under a vacuum.

Cx26 was resuspended with 10 μ l of 50 mM NH₄HCO₃, 5 % (v/v) acetonitrile and 2 M urea and incubated for 30 min at 37 °C. After dilution to approx. 0.5 M urea with 50 mM NH₄HCO₃ and 5 % (v/v) acetonitrile, Cx26 was digested overnight with 50–100 ng of Tryp, Chymo (both at 37 °C) or GluC (at 25 °C). Digestions were stopped with an equal volume of 10 % (v/v) methanol, 60 % (v/v) acetonitrile, 1 % (v/v) TFA and 0.1 % OG, and then dried under vacuum. Cx26 peptides were resuspended in 0.2 % TFA buffer compatible for desalting by micro-C₁₈ or micro-SCX ziptips. Alternatively, peptides from each digestion were recovered with 2.5 % (v/v) Poros-20R2 reversed-phase beads, as described above.

MS

Spectra were acquired using an ABI4800 mass spectrometer (Applied Biosystems) in a positive-ion mode with internal mass calibration using the Glu-fibrinogen peptide, ACTH₁₈₋₃₉ and Tryp autodigestion products (m/z 842.502 and 2211.096 a.m.u.) if present. MS spectra (m/z 800–3600) were processed by standard methods, including noise removal (2.0 S.D.), baseline correction and de-isotoping. Acquisition, processing and interpretation parameters were optimized further 'on-the-fly' to improve ionization and fragmentation of hydrophobic and lowabundance Cx26 peptides. The eight most and least intense ions per MALDI spot, with signal/noise ratios >25, were selected for subsequent MS/MS analysis in both 1 and 2 keV modes. MS/MS spectra were acquired using heavy CID (collision-induced dissociation) and higher gas pressure $(1 \times 10^{-6}$ compared with 1×10^{-7} Pa), laser intensity (for MS, 3200–3500; and for MS/MS, 3875 or higher), and 3125-5000 subspectra were collected to increase the possibility of hydrophobic peptide fragmentation and improve peptide signal intensity. Each spectrum was averaged with 3000 laser shots and smoothed with the Savitsky–Golay algorithm [FWHM (full width at half-maximum) = 9, polynomial order = 4].

GPS Explorer v3.5 (Applied Biosystems) was used to process MS and MS/MS spectra to submit the peak lists to the MASCOT v1.9 search engine (http://www.matrixscience.com; Matrix Science) for peptide identification against the MSDB v20060831, SwissProt v57.7 and NCBI (National Center for Biotechnology Information) v20090912 databases. Typically, the total ion score CI (confidence interval) was greater than 95%. The peptide sequences were additionally matched against a connexin-only database. For MS analyses, the precursor mass error tolerance was set to 20 p.p.m., whereas the MS/MS mass error tolerance was set to 0.3 Da, three missed cleavages were allowed, and carbamidomethyl [CAM-Cys (carboxyamidomethylcysteine)] and oxidation [Oxi-Met (methionine oxidation)] were set as variable modifications.

Other artifactual modifications potentially introduced during sample preparation, such as methylated glutamate, acetylated aspartate or C-terminus and formylated N-terminus, as well as contamination from human keratin, were excluded in peak lists used for database searches (FindPept; http://expasy.org/tools/ findpept.html). In separate database searches, to identify nontryptic and semi-tryptic cleavages in Tryp digestions, the enzyme parameter was set as 'none' or 'semi-Tryp'.

Peak lists were also matched against the FindMod (http:// au.expasy.org/tools/findmod/) and/or ProFound (http://prowl. rockefeller.edu) databases as 'unknowns' (parameters: rodentia 20–40 kDa, pI 0–14, mass tolerance 50 p.p.m.), and subsequently against the expected fragment masses of digested Cx26 to determine m/z peaks corresponding to modified Cx26 peptides. The Delta mass tables listing PTMs were also consulted (http://prowl.rockefeller.edu/aainfo/deltamassv2.html).

Parameters for individual Cx26 PTM analyses were set up separately in different searches and, in each case, the PTMs were set up as variable modifications. Suspected PTMs of Cx26 were rejected if the modified peptides were not found unmodified in the same digestion, there was lack of recurrence in different digestions, or when the error was greater than 20 p.p.m. Putative Cx26 PTMs in each spectrum were matched further by percentage CI or confirmed manually by peak assignment using Data Explorer v4.6 (Applied Biosystems).

RESULTS

Identification of peptides from Cx26 by MALDI–TOF/TOF-MS/MS

Enzymatic digestion and MS analysis of soluble proteins generally yields 40–60% sequence coverage [9,10]. Sequence coverage of membrane proteins is typically lower due to the hydrophobicity of transmembrane domains, which are resistant to proteolysis, and because the resulting hydrophobic peptides are difficult to extract from in-gel digestions, do not ionize well or are difficult to solubilize in buffers compatible with MS [9]. To address these problems, several preparative and analytic methods were applied to optimize Cx26 sequence coverage.

To maximize sequence coverage, in-gel and in-solution enzymatic digestions of Cx26 were carried out in parallel with Tryp, GluC and Chymo. After stepwise elution of peptides from the reversed-phase bead matrix, both MS and MS/MS spectra were acquired for peptides in each elution fraction. The matched m/z lists of Cx26 enzyme digestions are shown in Table 1. MS sequence coverage was 62.6% for Tryp (Table 1A), 7.9% for GluC (Table 1B) and 29.9% for Chymo (Table 1C). Total MS sequence coverage was 71.3%, with 29.9% of the Cx26 sequence (**A**)

Table 1 Sequence coverage of Cx26

Sequence coverage of Cx26 purified from HeLa cells by Tryp digestion was 62.6% (**A**), by GluC digestion was 7.9% (**B**) and by Chymo digestion was 29.9% (**C**). Overall MS sequence coverage was 71.3%, with 29.9% sequence confirmed by MS/MS (\checkmark). The cytoplasmic, membrane and extracellular domain boundaries in all of the Tables follow those described by Maeda et al. [15]. The modified residue is in bold, and the specific modification and amino acid position from the N-terminus of the peptide are given. CAM-Cys is the result of alkylation with 2-iodoacetamide.

| Commen by IVIS/IVIS | Calculated m/z | Observed m/z | Δ (p.p.m.) | Missed cuts | Peptide, observed modification | Domain |
|---------------------|----------------|--------------|-------------------|-------------|-------------------------------------------------------------------------------------------------------------------------|--------|
| | 1618.8257 | 1618.8197 | <u> </u> | 0 | M ¹ DWGTLQSILGGVNK ¹⁵ | NT |
| Ń | 1634.8207 | 1634.8186 | -1 | 0 | M ¹ DWGTLQSILGGVNK ¹⁵ , Oxi-Met at position 1 | NT |
| · | 2353.0222 | 2353.0203 | -1 | 0 | $E^{42}VWGDEQADFVCNTLQPGCK^{61}$, 2× CAM-Cys at positions 12 and 19 | E1 |
| \checkmark | 1820.8536 | 1820.8643 | 6 | 0 | N ⁶² VCYDHYFPISHIR ⁷⁵ , CAM-Cys at position 3 | E1/M2 |
| · | 2596.4507 | 2596.4451 | -2 | 0 | L ⁷⁶ WALQLIMVSTPALLVAMHVAYR ⁹⁸ | M2 |
| \checkmark | 1691.8856 | 1691.8792 | -3 | 2 | G ¹⁰⁹ EIKNEFKDIEEIK ¹²² | CL |
| · | 2049.0862 | 2049.0811 | -2 | 3 | G ¹⁰⁹ EIKNEFKDIEEIKTQK ¹²⁵ | CL |
| \checkmark | 1264.6425 | 1264.6384 | -3 | 1 | N ¹¹³ EFKDIEEIK ¹²² | CL |
| \checkmark | 1621.8431 | 1621.8359 | — 4 | 2 | N ¹¹³ EFKDIEEIKTQK ¹²⁵ | CL |
| · | 2863.3333 | 2863.2498 | -29 | 0 | V^{144} IFEAVF M YVFYI M YNGFF M QR ¹⁶⁵ , 3 \times Oxi-Met at positions 8,14 and 20 | M3/E2 |
| | 2452.0842 | 2452.0789 | -2 | 0 | C^{169} NAWPCPNTVDCFISRPTEK ¹⁸⁸ , 3× CAM-Cys at position 1, 6 and 12 | E2 |
| \checkmark | 1051.6735 | 1051.7043 | 29 | 1 | S ²²² KRPVLVPR ²³⁰ | CT/Tag |
| \sim | 836.5464 | 836.5432 | <u> </u> | 0 | R ²²⁴ PVLVPR ²³⁰ | CT/Tag |
| | 2866.2214 | 2866.2271 | 2 | 0 | G ²³¹ SYPYDVDPYALHNHNHNHNHN ²⁵⁴ | Tag |
| (B) | | | | | | |
| Confirmed by MS/MS | Calculated m/z | Observed m/z | Δ (p.p.m.) | Missed cuts | Peptide, modification | Domain |
| | 1167.6666 | 1167.6754 | 8 | 0 | K ¹⁰² KRKF M KGE ¹¹⁰ , Oxi-Met at position 6 | CL |
| | 1635.9363 | 1635.9386 | 1 | 1 | K ¹⁰² KRKFMKGEIKNE ¹¹⁴ | CL |
| | 1651.9312 | 1651.925 | <u> </u> | 1 | K ¹⁰² KRKF M KGEIKNE ¹¹⁴ , Oxi-Met at position 6 | CL |
| | 1114.6942 | 1114.6963 | 2 | 0 | I ¹²¹ KTQKVRIE ¹²⁹ | CL |
| (C) | | | | | | |
| Confirmed by MS/MS | Calculated m/z | Observed m/z | Δ (p.p.m.) | Missed cuts | Peptide, modification | Domain |
| | 2877.4637 | 2877.4177 | - 16 | 3 | R ³² IMILVVAAKEVWGDEQADFV C NTL ⁵⁶ , CAM-Cys at position 22 | M1/E1 |
| | 1233.5714 | 1233.5955 | 20 | 1 | C^{53} NTLQPGCKNV ⁶³ , 1 × CAM-Cys at position 1 or 8 | E1 |
| | 2447.1746 | 2447.1919 | 7 | 2 | Q^{57} PG C KNV C YDHYFPISHIRL ⁷⁶ , 1 × CAM-Cys at position 4 or 8 | E1/M2 |
| | 1859.0544 | 1859.0139 | -22 | 0 | R ⁹⁸ RHEKKRKF M KGEI ¹¹¹ . Oxi-Met at position 10 | CL |
| | 2543.1451 | 2543.1438 | -1 | 3 | F^{162} MQRLVKCNAWPCPNTVDC F^{181} , 3× CAM-Cys at positions 8, 13 and 19 | E2 |





intensity

2

% intensity

20

(1) MS spectrum of Cx26 Tryp digestion, with letters A–E indicating the peptides for which MS/MS spectra are subsequently shown. Inset, the exact location of these peptides in a schematic diagram of Cx26 showing the cytoplasmic, membrane (boxed) and extracellular domain boundaries as described by Maeda et al. [15]. (A) NT domain; (B) E1; (C) CL; (D) CT domain; (E) CT domain purification tag. Labelling of *b* and *y* ions in MS/MS spectra was according to Biemann et al. [14].

9.0

1923.0

confirmed by MS/MS sequencing (Figure 1), values significantly higher than typically observed in membrane protein analysis [10].

mass (m/z)

Sequences not covered by MS analyses were only from the transmembrane domains: a region of M1 proximal to the NT domain, a region of M3 proximal to CL and almost all of M4.

Figure 1 (panel 1) shows the MS spectra of one elution fraction from a Tryp digestion (showing approx. 32 % sequence coverage; peaks with m/z of approx. 1691, 2049 and 2353 a.m.u. are also from Cx26), and selected MS/MS spectra of peptides from each of the Cx26 domains: the NT domain (Figure 1A), E1 (Figure 1B), CL (Figure 1C), CT domain (Figure 1D) and the CT domain purification tag (Figure 1E).

Non-tryptic and semi-tryptic cleavages of peptides

(8(+1)

1216.2

mass (m/z)

1819.8

PYD(+1), DPY(+

612.6

To increase the possibility of hydrophobic peptide fragmentation and improve peptide ion signal intensities, MS and MS/MS spectra were acquired, as needed, using CID, higher gas pressure

2323.4

3027.0

Table 2 Peptides from non-tryptic and semi-tryptic cleavages of Cx26

Format designating modification and abbreviations are as in the legend to Table 1.

| Calculated m/z | Observed m/z | Δ (p.p.m.) | Peptide, modification | | |
|----------------|--------------|-------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------|-------|--|
| 1054.6118 | 1054.6056 | -6 | V^{27} LFIFRI M ³⁴ , Oxi-Met at position 8 | M1 | |
| 1136.5470 | 1136.5457 | -1 | N ¹¹³ EFKDIEEI ¹²¹ | CL | |
| 1168.5747 | 1168.5813 | 6 | A ¹⁴⁸ VF M YVFYI ¹⁵⁶ , Oxi-Met at position 4 | M3 | |
| 1433.6300 | 1433.6211 | -6 | L^{56} QPG C KNV C YDH ⁶⁷ , 1 × CAM-Cys at position 5 or 9 | E1 | |
| 1440.6543 | 1440.6287 | - 18 | P ²³⁴ YDVDPYALHNH ²⁴⁵ | Tag | |
| 1452.7151 | 1452.7059 | -6 | T ¹⁷⁷ VD C FISRPTEK ¹⁸⁸ , CAM-Cys at position 4 | E2 | |
| 1490.7308 | 1490.7432 | 8 | M ¹ DWGTLQSILGGVN ¹⁴ | NT | |
| 1552.6882 | 1552.6636 | 16 | D^{50} FV C NTLQPG C KN ⁶² , 2× CAM-Cys at positions 4 and 11 | E1 | |
| 1638.8639 | 1638.8557 | -5 | K ¹²⁵ VRIEGSLWWTYT ¹³⁷ | CL/M3 | |
| 1650.8019 | 1650.8164 | 9 | N ²⁰⁶ ITELCYLFIRYC ²¹⁸ | M4/CT | |
| 1656.8137 | 1656.8007 | - 8 | G ¹⁶⁰ FFMQRLVK C NAW ¹⁷² , CAM-Cys at position 10 | E2 | |
| 1666.808 | 1666.8086 | 0 | L^{210} CYLFIRYCSGKS ²²² , 2× CAM-Cys at positions 2 and 9 | M4/CT | |
| 1676.8036 | 1676.8248 | 12 | Y ¹⁵⁸ NGFFMQRLVK C N ¹⁷⁰ , CAM-Cys at position 12 | E2 | |
| 1697.8760 | 1697.8760 | 0 | 1 ¹⁴⁰ FFRVIFEAVF M Y ¹⁵² , Oxi-Met at position 12 | M3 | |
| 1784.9080 | 1784.9152 | 4 | S ¹³⁹ IFFRVIFEAVF M Y ¹⁵² , Oxi-Me at position 13 | M3 | |
| 1993.9921 | 1993.9855 | -3 | F ¹⁴¹ FRVIFEAVF M YVFY ⁻¹⁵⁵ , Oxi-Met at position 11 | M3 | |
| 2111.9094 | 2111.9385 | 14 | $F^{51}VCNTLQPGCKNVCYDH^{-67}$, 3× CAM-Cys at positions 3,10 and 14 | E1 | |
| 2185.0139 | 2185.0464 | 15 | F ¹⁶¹ FMQRLVKCNAWPCPNT ¹⁷⁷ , 2× CAM-Cys at positions 9 and 14, and Oxi-Met at position 3 | E2 | |
| 2364.0178 | 2364.03 | 5 | G ²³¹ SYPYDVDPYALHNHNHNHN ²⁵⁰ | Tag | |
| 2367.0745 | 2367.064 | <u> </u> | K ⁴¹ EVWGDEQADFVCNTLQPGCK ⁶¹ | E1 | |
| 2435.0916 | 2435.0681 | -10 | M^{157} YNGFFMQRLVKCNAWPCP ¹⁷⁵ , 2× CAM-Cys at positions 13 and 18, and 1× Oxi-Met at position 1 or 7 | E2 | |
| 2447.1746 | 2447.1985 | 10 | L^{56} QPG C KNV C YDHYFPISHIR ⁷⁵ , 1 × CAM-Cys at position 5 or 9 | E1/M2 | |
| 2447.1746 | 2447.1985 | 10 | Q^{57} PG C KNV C YDHYFPISHIRL ⁷⁶ , 1 × CAM-Cys at position 4 or 8 | E1/M2 | |
| 2451.1228 | 2451.1394 | 7 | I^{156} MYNGFFMQRLVKCNAWPC ¹⁷⁴ , 2× CAM-Cys at positions 14 and 19, and 1× Oxi-Met at position 2 or 8 | E2 | |
| 2501.2202 | 2501.2551 | 14 | P ¹⁷³ CPNTVDCFISRPTEKTVFTVF ¹⁹⁴ | E2/M4 | |
| 2510.1626 | 2510.1379 | -10 | K ¹⁶⁸ CNAWPCPNTVDCFISRPTEKT ¹⁸⁹ | E2 | |
| 2550.1938 | 2550.2397 | 18 | L^{166} VK C NAWP C PNTVD C FISRPTE ¹⁸⁷ , 1 × CAM-Cys at position 4, 9 or 15 | E2 | |
| 2566.2866 | 2566.292 | 2 | C^{202} ILLNITELCYLFIRYCSGKS ²²² , 2× CAM-Cys at two of positions 1, 10 or 17 | M4/CT | |
| 2566.3018 | 2566.292 | <u> </u> | I ¹²⁸ EGSLWWTYTTSIFFRVIFEA ¹⁴⁸ | M3 | |
| 2615.1196 | 2615.1445 | 10 | G ²³¹ SYPYDVDPYALHNHNHNHNHN ²⁵² | Tag | |
| 2705.1582 | 2705.1624 | 2 | A^{49} DFVCNTLQPGCKNVCYDHYFP ⁷⁰ , 3× CAM-Cys at positions 5, 12 and 16 | E1 | |
| 2894.2703 | 2894.2478 | - 8 | M ¹⁵⁷ YNGFFMQRLVK C NAWP C PNTVD C ¹⁸⁰ , 1 × CAM-Cys at position 13, 18 or 24 | E2 | |
| 3264.5427 | 3264.5261 | -5 | I^{145} FEAVF M YVFYI M YNGFF M QRLVK C ¹⁶⁹ , CAM-Cys at position 25, and 3× Oxi-Met at positions 7, 13 and 19 | M3/E2 | |
| 3227.4392 | 3227.468 | 9 | Y ¹⁵⁵ IMYNGFFMQRLVKCNAWPCPNTVDC ¹⁸⁰ , 2× CAM-Cys at two of positions 15, 20 or 26 | E2 | |

 $(1 \times 10^{-6} \text{ Pa})$ and laser intensity (>3875). Such changes can lead to better secondary fragmentation of peptides by in-source collisions. Additionally, in tryptic digestions some cleavages may not occur specifically at arginine or lysine residues due to residual chymotryptic activity, result from pseudotrypsin activity (Tryp autodigestion), partial enzymatic cleavage (i.e. one end of the peptide arises from non-specific enzymatic cleavage) or can occur by degradation during protein purification itself. Thus a number of other peptides were observed (Table 2), including those 'missing' from M1, M3 and M4, which were not predicted by an *in silico* digestion of Cx26. These were very low abundance in the spectra and not amenable to analysis that could reveal PTMs. However, if included, MS sequence coverage would rise to 92.6% of Cx26.

PTMs of Cx26

The PTMs of Cx26 identified by MS and by MS/MS analysis are listed in Table 3. All MS/MS sequence assignments that passed acceptance criteria after PTM database searches were manually verified, as most algorithms used for automated peptide identification are potentially less reliable in identifying spectra when the precursor ion is in the +1 charge state characteristic of MALDI. The peaks in each MS/MS spectrum were assigned as *b* and *y* ions [14], resulting, respectively, from

umber of missing' carboxyglutamation, methylation and phosphorylation.
Figure 2 shows two MS/MS spectra identifying PTMs.
These are the acetylation of the N-terminal methionine (Met¹)
(Figure 2A) and methylation of asparagine at position 254 (Asp²⁵⁴)
(Figure 2B). These PTMs correspond to mass additions of 42.01
and 14.01 a.m.u. respectively, to the amino acids.
Many more PTMs of Cx26 were indicated by MS spectra only. Their acceptance was subject to the stringent criteria listed

only. Their acceptance was subject to the stringent criteria listed above. These PTMs include multiple acetylations in the NT and CL domains, hydroxylation in the NT, CL and E2 domains, γ carboxylation of glutamate in the E1 and CL domains, methylation in the E1, M2 and CT domains, and phosphorylation in the CL and E2 domains, as well as PTMs (phosphorylation and methylation) in the CT domain purification tag. Representative MS spectra for these PTMs are shown in Figure 3.

cleavage of amide bonds from the N-terminal or C-terminal of the peptide. Where identification was by MS only, suspected

PTMs were rejected if the modified peptides were not found

unmodified in the same digestion, there was a lack of recurrence

of observation in different digestions or the error was >20

p.p.m. (see the Experimental section). Several putative PTMs

of Cx26 were identified, including acetylation, hydroxylation, γ -

All of the PTMs identified by MS or MS/MS are presented schematically in Figure 4. To determine whether the PTMs identified were at sites of $Cx26_{deaf}$, databases for mutations (http://davinci.crg.es/deafness and www.ncbi.nlm.nih.gov/omim)

Table 3 PTMs of Cx26

Suspected PTMs were rejected if (i) the modified peptides were not found unmodified in the same digestion, (ii) there was insufficient recurrence of observation in different digestions, (iii) the error was >20 p.p.m., or (iv) the amino acids involved were cysteine residues that were CAM-Cys-modified. Formats designating modifications are as in the legend to Table 1. Acetyl-Lys, acetylation of lysine; Acetyl-Met, acetylation of methionine; GGLU-Glu, glutamate γ -carboxylation; Hydroxyl-Asn, hydroxylation of asparagine; Methyl-Lys, methylation of lysine; Methyl-Argo, methylation of arginine; Phospho-Tyr, tyrosine phosphorylation; Phospho-Thr, threonine phosphorylation. Information is also shown schematically in Figure 4 (and its legend).

| Enzyme | Confirmed by MS/MS | Calculated <i>m/z</i> | Observed m/z | Δ (p.p.m.) | Missed cuts | Peptide, modification | Domain |
|--------|--------------------|-----------------------|--------------|-------------------|-------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------|
| Tryp | | 1650.8156 | 1650.8195 | 2 | 0 | M ¹ DWGTLQSILGGVNK ¹⁵ , Oxi-Met at position 1, and Hydroxyl-Asn at position 14 | NT |
| Tryp | | 1660.8363 | 1660.8247 | -7 | 0 | M ¹ DWGTLQSILGGVNK ¹⁵ , Acetyl-Met at position 1 | NT |
| Tryp | · | 1676.8312 | 1676.8179 | -8 | 0 | M ¹ DWGTLQSILGGVNK ¹⁵ , Oxi-Met at position 1, and Acetyl-Lys at position 15 | NT |
| Tryp | | 2367.0381 | 2367.064 | 11 | 0 | E ⁴² VWGDEQADFVCNTLQPGCK ⁶¹ , 2× CAM-Cys at positions 12 and 19, and Methyl-Lys at position 20 | E1 |
| Tryp | | 2383.9805 | 2383.9521 | - 12 | 0 | E ⁴² VWGDEQADFVCNTLQPGCK ⁸¹ , 2× CAM-Cys at positions 12 and 19, and 2× GGLU-Glu at positions 1 and 6 | E1 |
| Tryp | | 1777.8479 | 1777.8655 | 10 | 0 | N ⁶² VCYDHYFPISHI R ⁷⁵ , Methyl-Arg at position 14 | E1/M2 |
| Tryp | \checkmark | 1834.8694 | 1834.8735 | 2 | 0 | N ⁶² VCYDHYFPISHIR ⁷⁵ , CAM-Cys at position 3 and Methyl-Arg at position 14 | E1/M2 |
| Chymo | | 1602.8976 | 1602.8877 | -6 | 1 | A ⁹⁶ YRRHE KK RK ^{F106} , 2× AcetyI-Lys at two of positions 7, 8 or 10 | M2/CL |
| GluC | | 2252.1743 | 2252.1506 | - 11 | 2 | K ¹⁰² KRKFMKGEIKNEFKD ⁻¹¹⁷ , Oxi-Met at position 6, and 5× AcetyI-Lys at five of positions 1, 2, 4, 7, 11 or 15 | CL |
| Chymo | | 909.4676 | 909.4496 | - 20 | 0 | K ¹¹² NEFKDI ¹¹⁸ , Hydroxy-Asn at position 2 | CL |
| Chymo | | 937.4625 | 937.4501 | — 13 | 1 | K ¹¹² NEFKDI ¹¹⁸ , GGLU-Glu at position 3 | CL |
| GluC | | 1565.8298 | 1565.8145 | — 10 | 2 | I ¹¹⁸ EEIK T QKVRIE ¹²⁹ , Phospho-Thr at position 6 | CL |
| Tryp | | 2694.2837 | 2694.2505 | - 12 | 2 | L ¹⁶⁶ VK CN AWP CPN TVD C FISRPTEK ¹⁸⁸ , 1 × CAM-Cys at position 4, 9 or 15, and 1 × Hydroxy-Asn at position 5 or 11 | E2 |
| Tryp | | 3032.2307 | 3032.2522 | 7 | 1 | L ¹⁶⁶ VKCNAWPCPNTVDCFISRPTEK ¹⁸⁸ , 3× CAM-Cys at positions 4, 9 and 15, and 3× Phospho-Thr at positions 12, 18 and 21 | E2 |
| Tryp | | 843.4028 | 843.3899 | — 15 | 1 | Y ²¹⁷ CSGKSK ²²³ , CAM-Cys at position 2, and 1 × Methyl-I vs at position 5 or 7 | СТ |
| Tryp | \checkmark | 2880.2375 | 2880.2437 | 2 | 0 | G ²³¹ SYPYDVDPYALHNHNHNHNHNHNHN ²⁵⁴ , 1× Methyl-Asn at position 22 | Tag |
| Tryp | | 2946.1877 | 2946.1924 | 2 | 0 | $G^{231}SYPYDVDPYALHNHNHNHNHNHN^{254}$, 1 × Phospho-Tyr at position 3, 5 or 10 | Tag |

were consulted. A significant fraction of the PTMs identified by MS and MS/MS are at $Cx26_{deaf}$ sites (Table 4). These were hydroxylation at Asn¹⁴, γ -carboxylation at Glu⁴² and Glu⁴⁷, methylation at Arg⁷⁵, acetylation at Lys¹⁵ and Lys¹⁰², and phosphorylation at Thr¹²³ and Ser¹⁸³.

DISCUSSION

PTMs of proteins are diverse and play important modulatory roles in trafficking, biogenesis and function [8]. In the present study, MALDI-TOF/TOF-MS was employed to identify PTMs of Cx26 expressed in HeLa cells. Protocols for sample preparation for MS and MS/MS analysis were optimized to improve sequence coverage, ionization and fragmentation of hydrophobic regions of the protein, since they present difficulties for MS analysis. These refinements included the use of highly pure connexin, parallel in-gel and in-solution digestion by three proteases with distinct cleavage specificities, peptide fractionation with a reversedphase matrix, and adjustment of the MS/MS and collision parameters to increase fragmentation. This resulted in 71.3% confirmed sequence coverage by MS and 29.9% confirmed by MS/MS sequencing. Low-abundance peptides, not amenable to MS/MS analysis and arising from other sources, such as insource collisions, partial enzymatic cleavage, pseudotrypsin or chymotryptic activity in tryptic digestions, were observed. If these

are included, MS sequence coverage was 92.6%. The results indicate a variety of PTMs of Cx26 distributed over the protein sequence, several of which are at sites that when mutated in Cx26 (and other connexins) give rise to human pathologies.

Analysis of Cx26 PTMs using MALDI-MS/MS

According to the recent crystal structure of Cx26 [15], approx. 47% of the sequence is in transmembrane regions. This high proportion of hydrophobic sequence presents difficulties for digestion by proteases and subsequent peptide analysis by MS, as indicated by the absence in Table 1 of peptides from short regions of M1 and M3, and the entire M4 domain. The two transmembrane helices assigned in the crystal as forming the outer wall of the pore (i.e. regions exposed to lipid for almost all of their lengths that are within the membrane) are part of M3 and principally M4, whereas M1, the principle pore-lining helix, is atypically hydrophobic [using ProtParam (http://au.expasy.org/tools/protparam.html), the average hydropathy of M1 is 1.85 compared with 1.20, 1.05 and 1.31 for M2, M3 and M4 respectively]; M1 does contain amino acids involved in hydrophobic intra-protomer interactions.

MALDI-TOF-MS typically has a detection limit of low femtomole to high attomole, but for membrane protein and peptides with hydrophobic character, the detection limit is much higher and signal intensity (peptide ionization) is usually

Table 4 PTMs at sites of disease-causing mutations

This Table summarizes information regarding the disease-causing mutations of Cx26 that are at sites of PTMs identified in the present study. Abbreviations: n, non-syndromic; s, syndromic; r, recessive; d, dominant. Information is also shown schematically (see Figure 4 and its legend).

| Mutation (type) | Domain | Mechanism (if known) | Observed PTM |
|-----------------------------------------------------------------|--------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------|
| Asn ¹⁴ Tyr (s, d*) and Asn ¹⁴ Lys (s, d†) | NT | Asn ¹⁴ Tyr: Keratitis-Ichthyosis-Deafness syndrome*, forms functional junctional channels, reduced junctional conductance, change to local structural flexibility of the NT domain [30]; Asn ¹⁴ Lys: Clouston syndrome†, caused one hemichannels and loss of voltane-dependence [51 52] | Hydroxylation |
| Lvs ¹⁵ Thr (n. r) | NT | No information on functional effect [53]. | Acetvlation |
| del-Glu ⁴² (s, d) | E1 | Forms junctional plaques but not functional junctional channels, behaves as dominant-negative with Cx26, Cx30 and Cx43 [35,36]. | γ -Carboxyglutamation |
| Glu ⁴⁷ Lys (n, r) | E1 | Normal trafficking, but does not form functional hemichannels or junctional channels [37]. | γ -Carboxyglutamation |
| $Arg^{75}Gin$ (n, d) and $Arg^{75}Trp$ (s, d‡) | E1 | Forms junctional plaques in some reports, not in others, but no functional junctional channels, forms hemichannels with altered voltage-dependence and reduced permeability, behaves as dominant-negative with Cx26, Cx30 and Cx43 [42,54,55]. (Arq ⁷⁵ Trp for Palmoplantar keratoderma† [55]). | Methylation |
| del-Lys ¹⁰² (n, r) | CL | No information on functional effects [56]. | Acetylation |
| Thr ¹²³ Asn (n, d§) | CL | No information on functional effects [57,58]. | Phosphorylation |
| Ser ¹⁸³ Phe (s, d‡) | E2 | Behaves as dominant-negative with Cx26, impaired formation of junctional channels with those forming being Lucifer yellow permeable [48]. | Phosphorylation |

* http://www.ncbi.nlm.nih.gov/entrez/dispomim.cgi?id=148210 † http://www.ncbi.nlm.nih.gov/entrez/dispomim.cgi?id=129500

‡ http://www.ncbi.nlm.nih.gov/entrez/dispomim.cgi?id=148350

 \S possible Cx26_{deaf} mutation (http://davinci.crg.es/deafness)



Figure 2 MS and MS/MS spectra showing PTMs of Cx26

All modified peptides were also found in their unmodified state in the same digestions. (**A**) MS spectrum of the Cx26 Tryp digestion (lower panel) and MS/MS spectrum of the NT domain showing N-terminal acetylation (A) of methionine (position Met¹). (**B**) MS spectrum of Cx26 Tryp digestion (lower panel) and MS/MS spectrum of the CT domain purification tag showing N-terminal methylation (**M**) of asparagine (position Asp²⁵⁴). See Table 1 for full details of the sequence coverage, and Table 3 for other PTMs. Labelling of *b* and *y* ions in MS/MS spectra was according to Biemann et al. [14].





Figure 3 MS spectra suggesting PTMs of Cx26 for which high-quality MS/MS spectra were not obtained

Representative MS spectra for the indicated modifications. See the Experimental section and the legend to Table 3 for acceptance criteria. (**A**) Acetylation (A) in the CL proximal to the second transmembrane domain. (**B**) hydroxylation (H) in the NT domain proximal to the first transmembrane domain. (**C**) glutamate γ -carboxylation (G) in the central part of the CL. (**D**) phosphorylation (P) of the CT domain purification tag. (**E**) phosphorylation (P) in the central part of the CL. Amino acids marked with asterisks (*) are sites of Cx26_{deal} mutation. See Table 1 for details of sequence coverage and Table 3 for other PTMs.

much lower than for water-soluble peptides. This may be a consequence of membrane proteins having a greater tendency than soluble proteins to remain folded in the gas phase; additionally, MALDI-TOF/TOF-MS/MS sequencing using CID depends on protonation of amide groups, and folded hydrophobic regions are poorly ionized. With our optimized MALDI parameters, the detection sensitivity was improved but was still insufficient to detect low-abundance PTMs, resulting in incomplete coverage by MS/MS sequencing.

In theory, ESI-LC-QTOF-MS [electrospray-coupled LC with QTOF (quadrupole TOF)-MS] would provide improved and/or complementary ionization for MS/MS sequencing of peptides with hydrophobic character, since peptides can carry multiple charges, rather than single charges (here, +1) as in MALDI–TOF-MS. Furthermore, this technology might allow more sensitive detection of, in particular, sites of phosphorylation of Cx26. The dominant neutral loss of phosphoric acid from phosphorylated peptides when using MALDI and a TOF



Figure 4 Schematic representation of the identified PTMs of Cx26

Each amino acid of Cx26, and of the tandem CT domain epitopes used for protein purification, is shown individually. Filled black rectangles indicate regions for which MS/MS confirmation was achieved, with sequences for which only MS data were available is shown by filled grey rectangles. If rectangles are not filled (i.e. white), then no sequence information/peptides were obtained by MALDI–TOF (see Tables 1 and 2 for peptide lists and the text for discussion of their absence). The PTMs (see Table 3 and 4, and their legends) are identified using the following abbreviations: acetylation (A), hydroxylation (H), γ -carboxyglutamation (G), methylation (M) and phosphorylation (P). Sites of Cx26_{deat} disease-causing mutation, where PTMs were identified, are designated by single-letter amino acid code, are oval-shaped and marked inset by an asterisk (*). The positions of the cytoplasmic, membrane (stippled lines) and extracellular domain boundaries are as described by Maeda et al. [15].

analyser in reflector mode ('post-source decay') typically results in insufficient fragmentation of such phosphopeptides during MALDI–TOF/TOF-MS/MS. However, even by ESI-LC-QTOF-MS there may still be some element of uncertainty for assignment, especially where there is more than one potential location in the peptide for PTM. In this case, confirmation requires MS³ of product–phosphoric acid ions using an 'ion trap', which is not possible with the MALDI–TOF/TOF-MS employed in the present study. However, ESI-LC-QTOF-MS suffers from other technical challenges, notably being less tolerant than MALDI–TOF/MS to non-peptide contaminants such as salt, and especially lipid or detergent that may be likely to remain bound post-digestion to hydrophobic peptides.

One previous investigation of connexin PTMs by proteomic methods (Cx44 and Cx49) utilized HPLC for desalting, isolation and concentration of peptides, a custom-built MALDI-QTOF system, custom peptide identification software, two proteases and in-solution digestion, but achieved substantially less MS and MS/MS sequence coverage than in the present study, and allowed a greater margin of error for assigning PTMs [16]. Aside from N-terminal acetylation, several sites of phosphorylation were found in the CL and long CT domains of these connexins. The sites were identified, as outlined above, by neutral loss of phosphoric acid from MS/MS fragment ions. Peptides were not recovered

from the transmembrane and extracellular domains, the proximal portions of the CT domain, or from substantial portions of the NT domain (Cx44) or the CL domain (Cx49).

The greater sequence coverage in Cx26 MS and MS/MS sequencing in the present study can be attributed to the various technical improvements in MALDI–TOF-MS and MALDI–TOF/TOF-MS/MS spectra acquisition described above and to the purity of the starting material (in the study of Cx44 and Cx49, only 10–18 out of 600–900 MS peaks were from the connexins). The purification of Cx26 employed in the present study used tandem CT domain tags and yielded a protein of high purity. This contributes to greater sensitivity in peptide and PTM detection, since non-connexin/contaminating peptides absorb 'energy' and suppress ionization of peptides of interest. Previous work using less optimized purifications and MS running conditions identified only a small number of PTMs of Cx26, which were also found in the present study (hydroxylation at Asn¹⁴ and γ -carboxylation at Glu¹¹⁴ [13]).

PTMs of Cx26

Several different PTMs of Cx26 were identified, including acetylation, hydroxylation, γ -carboxyglutamation, methylation

and phosphorylation. The PTMs confirmed by MALDI–TOF/ TOF-MS/MS sequencing can be viewed with high confidence. The others were identified by mass consistency and accuracy of expected peptide fragments. The limitations of using single m/z peaks to identify PTMs are acknowledged; however, strict criteria were applied to minimize the chance of erroneous PTM identification (see the Experimental section). In particular, assignment of the PTMs was made using criteria that each was supported by the observation of re-occurrence of the modified peptide in different digestions along with identification of the unmodified peptide in the same digestion. The PTMs identified from MS by these criteria are highly likely, but not certain, to be correct. For simplicity, below, PTMs may be referred to as 'identified', although this is not, in the strictest sense, true for those resulting from MS analysis alone.

Acetylation of Cx26

Acetylation was identified at the N-terminal methionine (Met¹, by MS/MS), elsewhere in the NT domain and at several lysine residues in the CT domain (Lys¹⁵, two among Lys¹⁰², Lys¹⁰³ and Lys¹⁰⁵, and all of Lys¹⁰⁸, Lys¹¹² and Lys¹¹⁶; by MS). N-terminal acetylation is a common irreversible cotranslational modification. Proteins with the Met¹-Asp² sequence, such as Cx26, undergo an obligatory N-terminal acetylation. This can affect a variety of factors, most prominently protein stability due to the blocked N-terminus, but also protein function and subcellular targeting [17].

Acetylation of the N-terminus eliminates its positive charge. In connexins, the NT domain folds into the aqueous pore to act as a voltage sensor and gate [15,18–20]. Thus charge elimination at the N-terminus may have a profound impact on channel gating. It was reported that the N-terminal methionine residue in Cx44 and Cx49 is endogenously removed and the newly exposed N-terminal glycine residue becomes acetylated [16], so N-terminal acetylation may be a widespread connexin PTM.

Unlike N-terminal acetylation, acetylation of internal lysine residues is post-translational and reversible. An internal lysine acetylation in the NT domain (Lys¹⁵) was identified, as were several other lysine residue acetylations in a short segment of the CL domain proximal to M2. Internal lysine residue acetylation can regulate diverse functions, including protein–protein interaction and protein stability [21]. Lysine residue acetylation can also affect other PTMs, e.g. lysine residues are targets for ubiquitination and acetylation can block this modification [22].

According to the Cx26 crystal structure, Lys¹⁵ is at the outer edge of the cytoplasmic entrance/vestibule of the channel pore, so charge removal by acetylation could affect charge selectivity to permeants. Perhaps more important, Lys¹⁵ is near the 'hinge' region at which the NT domain turns into the pore to form the voltage sensor/gate [15,18], so a structural or charge modification at this site could affect voltage gating as well.

Lys¹⁵Thr is a Cx26_{deaf} mutation. The acetylation and this mutation both remove the positive charge. In Cx32, this position is an arginine residue, and the Arg¹⁵Gln mutation, which also eliminates positive charge, causes a peripheral neuropathy. Interestingly, Arg¹⁵Gln channels are functional, but have altered voltage sensitivity [23,24], consistent with the structural inferences. There is no functional information about Cx26_{deaf} Lys¹⁵Thr channel function, but, extrapolating from Cx32 Arg¹⁵Gln, one could propose that acetylation/charge removal at this site in Cx26 has a similar effect on channel voltage sensitivity, which results in non-syndromic deafness.

Several internal lysine residues are potentially acetylated in the Cx26 CL near M2. The connexin CL is involved in gating by pH and related intramolecular interactions [25,26], so dynamic modulation of the charge of this domain by multiple acetylations could modulate these processes, as occurs in other proteins [22].

In light of this, it is notable that deletion of Lys¹⁰² is a Cx26_{deaf} mutation. Acetylation or del-Lys¹⁰² eliminates a negative charge at this position. There is no functional information about Cx26_{deaf} del-Lys¹⁰²; however, in the CL of Cx32, Glu¹⁰²Gly eliminates the positive charge, as would acetylation, alters pH sensitivity and causes a mild neuropathy [27]. Thus it appears that a lack of charge at this position has a pathological effect on pH gating for both connexins.

Hydroxylation of Cx26

Hydroxylation was identified at Asn^{14} in the NT domain, at Asn^{113} in the CL, and at either Asn^{170} or Asn^{176} in E2. Protein hydroxylation is an irreversible modification that occurs at aromatic residues in a wide variety of proteins, but is less common at asparagine residues. Hydroxylated asparagine is found in several proteins, such as hypoxia-inducible transcription factors, epidermal growth factor and vitamin K clotting factors, where they are associated with high-affinity Ca²⁺-binding sites [28]. Cx26 does not contain the putative consensus sequence for hydroxylation by 2-oxoglutarate-dependent dioxygenases, but it presumes a characteristic β -fold [29].

There are two Cx26_{deaf} mutations at Asn¹⁴: Asn¹⁴Tyr and Asn¹⁴Lys. Asn¹⁴ is at the 'hinge' region that is, as noted above, where the NT domain folds into the pore [15,18], so its modification would be expected to affect (voltage) gating. Indeed, the Asn¹⁴Tyr mutation alters local peptide flexibility [30] and leads to reduced junctional conductance, consistent with altered gating, whereas the introduction of positive charge by Asn¹⁴Lys mutation generates open hemichannels, as well as functional gap junction channels, but they lack voltage-dependence. Asn¹⁴Tyr does not introduce a charge, so could be considered the mutation that better approximates the lack of hydroxylation.

Asn¹⁷⁶ is one of the residues in the Cx26 crystal structure suggested to be directly involved in the docking interaction between apposed hemichannels [15]. The Asn¹⁷⁵Asp mutation in Cx32 (corresponding to Asn¹⁷⁶ in Cx26) causes a peripheral neuropathy, but the effect on channel function has not been characterized [31].

Glutamate γ -carboxylation of Cx26

Glutamate γ -carboxylation was identified at Glu⁴² and Glu⁴⁷ in E1, and at Glu¹¹⁴ in the CL. This irreversible modification generates high-affinity Ca²⁺-binding sites; addition of the negatively charged carboxy group forms a Ca²⁺-chelation site with the existing glutamate side-chain carboxy group [8,32]. In this context, the position of these modifications in E1 is intriguing as, according to the crystal structure of Cx26 [15], they are in a 3₁₀ helix involved in Ca²⁺-sensitive 'loop gating' of connexin hemichannels, i.e. the gate that keeps unapposed hemichannels closed [33,34].

Intriguingly, both Glu^{42} and Glu^{47} are sites of $Cx26_{deaf}$ mutations. Deletion of Glu^{42} or $Glu^{47}Lys$ allows formation of junctional channels, but they are non-functional [35–37], perhaps indicating a loop gate malfunction, e.g. remaining closed in spite of hemichannel docking to form junctional channels. Other studies have identified Glu^{42} as a major determinant of the charge selectivity of the channel [38]. The $Cx26_{deaf}$ $Glu^{47}Lys$ mutation

eliminates the possibility of γ -carboxyglutamation, but imposes the same charge that γ -carboxyglutamation would, suggesting that it is the PTM itself and not the charge alteration itself that is the key factor for normal channel function.

Glutamate γ -carboxylation was identified in the CL, which contains a high density of glutamate resides (four between residues 114 and 120), although only one was detected so modified in the present study. As γ -carboxylated glutamate residues can mediate strong Ca²⁺-dependent interactions with membranes [32], these could mediate Cx26 sensitivity to cytosolic Ca²⁺ (proposed previously for Cx26 [13]). The Glu¹¹⁴Gly mutation apparently has no dramatic effect on dye permeability (pH sensitivity untested) [39]. Introduction of an increased negative charge into the CL may also modulate the various gating processes and related intramolecular interactions involving it, as mentioned above.

Methylation of Cx26

Methylation was identified at Lys⁶¹ in E1, at Arg⁷⁵ at the E1/M2 border (by MS/MS) and at either Lys²²¹ or Lys²²³ in the CT domain. Methylation at arginine and lysine residues is relatively widespread, irreversible and implicated in a wide spectrum of cellular processes, including protein stability, activity and protein–protein interactions [40]. In most cases, its primary effect follows from increased side-chain bulk and hydrophobicity.

The most intriguing methylation identified is at Arg^{75} , a site of two $Cx26_{deaf}$ mutations that inhibit junctional coupling in Cx26 as well as Cx32 [41,42]. Arg^{75} Trp, unlike Arg^{75} Gln, forms somewhat functional hemichannels [43]. These mutations both eliminate the positive charge, as well as the possibility of methylation, so the consequences of methylation itself cannot be inferred. However, it should be noted that Arg^{75} is at the E1/M2 border (i.e. the aqueous–lipid interface) and so its increased side-chain bulk and/or hydrophobicity after methylation may influence the positioning of M2 and E1 relative to the membrane.

In Cx32, the site that corresponds to Cx26 Lys²²¹ is Arg²²⁰, which conserves the charge and the potential for methylation. The Arg²²⁰Gly mutation in Cx32 forms junctional plaques and causes neuropathy, but the function of the junctional channels has not been assessed [41].

Phosphorylation of Cx26

Phosphorylation was identified at Thr¹²³ in the CL, at Thr¹⁷⁷ in E2, at Ser¹⁸³ and Thr¹⁸⁶ near the E2/M4 border, and at one of Tyr²³³, Tyr²³⁵ or Tyr²⁴⁰ in the epitope tag. To date, Cx26 has not been definitively reported to be a phosphoprotein. Our present findings indicate potential phosphorylation in the CL, a domain in which at least two other connexins (Cx36 and Cx56) are phosphorylated [44,45]. In contrast with CT domain phosphorylation, the effects of CL phosphorylation are underexplored. Thr¹²³ is a possible Cx26_{deaf} mutation, but there is no information on the functional effects of mutation at this site.

Several phosphorylation sites are indicated in E2. Such modifications could occur by intracellular kinase activity, but could also be introduced by ecto-kinases found on many cells, including HeLa cells, which modulate other membrane proteins [46, 47]. According to the Cx26 crystal structure [15], one of the suspected sites in E2 (Thr¹⁷⁷) forms a docking interaction with the apposed hemichannel.

Interestingly, the $Cx26_{deaf}$ Ser¹⁸³Phe mutation significantly reduces the number of junctional channels formed [48], but those formed retain normal dye permeability. It is therefore unknown if the primary defect is in trafficking, junctional formation or greater internalization of undocked hemichannels. The corresponding Ser¹⁸²Thr mutation in Cx32 that causes neuropathy also results in non-functional junctional channels [24]. Given the biological reversibility of phosphorylation, this might imply a role for it in regulating hemichannel docking at this site in both connexins.

Phosphorylation (and methylation; by MS/MS) was also indicated in the HA epitope, which may be relevant to evaluation of the function or cell biology of connexins carrying this tag.

Cx26 PTMs and deafness mutations

Mutations in the Cx26 gene (*GJB2*) account for over 50% of autosomal recessive non-syndromic congenital deafness, the most common form of genetic deafness, and 10-30% of sporadic cases. More than 50 mutations in the coding region of *GJB2* have been reported to be associated with autosomal-recessive (*DFNB1*) and autosomal-dominant (*DFNA3*) forms of hearing impairment [2,3].

The mechanism(s) by which deafness is caused by mutations that delete or alter the function of Cx26 is unknown. It has been proposed that disruption of intercellular K⁺ recycling in the cochlea is involved [5], leading to reversed operation of a glutamate transporter and accumulation of extracellular glutamate, which may inhibit synthesis of glutathione, causing death from oxidative stress [4].

Previous evidence has suggested that disruption of intercellular molecular signalling, rather than K^+ recycling, is the important factor. Some Cx26_{deaf} mutations produce channels with unaltered K^+ permeability, but with disrupted molecular permeability [49,50]. In addition, when Cx26 is genetically deleted, the remaining Cx30 junctional channels, found in the same locations, are highly permeable to K^+ [4]. Thus changes in the character, degree or regulation of connexin channel molecular permeability, as opposed to ionic conductance, appear to be sufficient to cause deafness and/or skin pathology.

The location of PTMs identified in the present study at which there are known deafness-causing mutations are indicated in Figure 4. Information about the functional effects of mutation at these sites is summarized in Table 4. Of the eight $Cx26_{deaf}$ sites at locations of PTMs, five are dominant mutations and four have syndromic phenotypes.

At this point it is difficult to make mechanistic links between the presence of specific PTMs normally at these sites and disruptions of normal Cx26 function when these sites have $Cx26_{deaf}$ mutations. Each $Cx26_{deaf}$ mutation alters the sequence, which can have its own effects, in addition to eliminating the (proper) PTM. It is unknown whether the detected PTMs are introduced as part of normal channel maturation, degradation or modulatory processes. However, if a particular PTM is part of a degradatory process and does not occur due to mutation at the site, one would expect the result to be increased channel function rather than decreased or compromised function. Such an effect is not observed, so one can presume that the observed irreversible PTMs at Cx26_{deaf} sites are to promote proper channel maturation and function, rather than proper degradation. Of the observed PTMs, only phosphorylation and acetylation are reversible, so one cannot infer a priori whether their presence or absence favours channel function.

It is notable that most of the identified PTMs, and all of those at $Cx26_{deaf}$ locations (with the possible exception of Ser¹⁸³, noted above), are in regions known or strongly suspected from biophysical work and/or the recent Cx26 crystal structure to be crucial for proper channel function: Asn¹⁴ and Lys¹⁵ are involved in voltage gating, Glu⁴² and Glu⁴⁷ are involved in loop gating and Arg⁷⁵ interacts directly with residues in M1, the pore-lining helix.

Although there is little functional information about specific PTM sites in the CL loop (Lys¹⁰², Asn¹¹³ and Glu¹¹⁴), it is recognized that the CL is a primary site of connexin channel functional modulation. One would anticipate that in most cases lack of proper PTMs would substantially affect trafficking, but the available evidence does not indicate this; many other $Cx26_{deaf}$ mutations do affect trafficking or channel assembly, but these were not identified in the present study as sites of PTM. Of the five PTM sites that are also sites of $Cx26_{deaf}$ mutation for which there are relevant data, all form junctional channels and, of those, three form functional junctional channels. Thus a case can be made that disease-causing disruption of PTMs are more likely to affect mature channel function than biogenesis.

Several of the disease mutations that do form functional channels are at sites at which PTMs have been identified. These are particularly interesting targets for future investigation.

Conclusions

The overall high sequence coverage of Cx26 achieved using MALDI–TOF and MS/MS made it possible to identify several new sites for PTM of Cx26. Many of the sites of PTM are at sites of Cx26 disease-causing mutation. Mutation at these sites could preclude these PTMs from playing important roles in connexin channel biogenesis and/or function. Such PTMs are promising targets to study the regulatory functions of connexin channel gating and permeation properties.

AUTHOR CONTRIBUTION

Darren Locke conceived and designed the experiments; Shengjie Bian and Darren Locke performed the experiments; Shengjie Bian, Andrew Harris and Darren Locke analysed the data; Hong Li contributed reagents/materials/analysis tools; Andrew Harris and Darren Locke wrote the paper.

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