Peripheral nerve pathology in sickle cell disease mice
Katelyn E. Sadler, Tylor R. Lewis, Tyler B. Waltz, Joseph C. Besharse, Cheryl L. Stucky

Abstract

Introduction: Many patients with sickle cell disease (SCD) suffer from chronic pain, which is often described as neuropathic in nature. Although vascular and inflammatory pathology undoubtedly contribute to the SCD pain experience, the nociceptive signals that ultimately drive symptoms are detected and transmitted by peripheral sensory neurons. To date, no systematic histological examination of peripheral nerves has been completed in patients or mouse models of SCD to diagnose disease-related neuropathy.

Objectives: In this brief report, we compared peripheral nerve morphology in tissues obtained from Berkeley transgenic SCD mice and control animals.

Methods: Sciatic nerves were visualized using light and transmission electron microscopy. Myelin basic protein expression was assessed through Western blot. Blood–nerve barrier permeability was measured using Evan’s blue plasma extravasation.

Results: Peripheral fibers from SCD mice have thinner myelin sheaths than control mice and widespread myelin instability as evidenced by myelin sheath infolding and unwrapping. Deficits are also observed in nonmyelinating Schwann cell structures; Remak bundles from SCD nerves contain fewer C fibers, some of which are not fully ensheathed by the corresponding Schwann cell. Increased blood–nerve barrier permeability and expression of myelin basic protein are noted in SCD tissue.

Conclusions: These data are the first to characterize Berkeley SCD mice as a naturally occurring model of peripheral neuropathy. Widespread myelin instability is observed in nerves from SCD mice. This pathology may be explained by increased permeability of the blood–nerve barrier and, thus, increased access to circulating demyelinating agents at the level of primary sensory afferents.

Keywords: Sickle cell disease, Myelinopathy, Sciatic nerve

1. Introduction

Sickle cell disease (SCD) is a hemoglobinopathy associated with many neurological complications including stroke, seizure, and pain. While pathologically diverse, all these symptoms are ultimately mediated by neurons, the fundamental unit of the nervous system. Despite the collective impact that these neurological symptoms have on patient quality of life, no studies have systematically examined the basic histological properties of neural tissue isolated from either patients with SCD or transgenic SCD mouse models to assess disease-related pathology.

In this report, sciatic nerves from Berkeley transgenic SCD mice (Berk SS) were examined using light and transmission electron microscopy (TEM). The sciatic nerve was chosen for these studies because of its large size and involvement in peripheral somatosensation. Dysregulated sensory systems lead to intense pain which is the leading cause of emergency department visits for patients with SCD. In both mouse and human, the sciatic nerve contains the thickly myelinated axons of motor efferents and the thinly myelinated and unmyelinated axons of sensory afferents, many of which are nociceptors. Here, we report skewed myelinated axon diameter distributions and decreased Remak bundle C-fiber density in Berk SS nerves. We also report decreased myelin sheath thickness and aberrant myelin sheath structures in nerves isolated from Berk SS mice. These changes are accompanied by an increase in myelin basic protein (MBP) expression and blood–nerve barrier permeability. These data are the first demonstration of myelin and axonal changes in SCD neural tissues, a phenotype that may contribute to many common SCD complications.

2. Methods

2.1. Mouse model and tissue collection

Berkeley transgenic mice homozygous for the sickle β globin gene were compared with control B6; 129 mice. Sciatic nerves (10 mm length isolated ~5 mm distal to hip) were obtained from...
male and female mice aged 8 to 12 weeks. All protocols were in accordance with National Institutes of Health guidelines and approved by the Institutional Animal Care and Use Committee at the Medical College of Wisconsin.

2.2. Light and transmission electron microscopy

Freshly dissected sciatic nerves were prepared as previously described. For light microscopy, semithin sections (0.5 μm) stained with toluidine blue were imaged on a Nikon Eclipse TE300 microscope. For TEM, thin sections (70 nm) were stained with uranyl acetate and lead citrate, then imaged on a Hitachi H-600 transmission electron microscope. Image analysis was completed using ImageJ software.

2.3. Myelin basic protein Western blot

Sciatic nerves were snap frozen on dry ice, pooled for individual animals then homogenized in lysis buffer (50 mM Tris base pH 8, 0.5% NP-40, 10% glycerol, 0.1 mM EDTA, 250 mM NaCl, 1X Halt protease inhibitor cocktail). Total protein content was analyzed using a BCA Protein Assay Kit (Pierce; ThermoFisher, Waltham, MA); 20 μg of protein was separated on 4% to 12% Bis-Tris gradient gels and transferred to nitrocellulose membranes. Membranes were blocked in Odyssey blocking buffer for 1 hour and incubated with mouse anti-MBP (abcam #ab62631, 1:1000; Abcam, Cambridge, United Kingdom) and rabbit anti-β-tubulin (abcam #ab6046, 1:10,000) primary antibodies for 1 hour. Blots were washed with 0.1% Tween-20 in PBS and incubated with donkey anti-rabbit 800 (LI-COR #926-32213, 1:15,000) and donkey anti-mouse 880 (LI-COR #926-68072, 1:15,000) secondary antibodies for 1 hour. Blots were scanned on an Odyssey infrared imaging system, and band densitometry was assessed using Image Studio Lite software. For each sample, MBP (23 and 18 kDa bands) was normalized to β-tubulin; SCD samples were then normalized to control average.

2.4. Evan’s blue extravasation

Evan’s blue (50 mg/mL stock, 80 mg/g body weight) was injected into the tail vein. Fifteen minutes after injection, mice were sacrificed and both sciatic nerves were harvested, weighed, and snap frozen. Nerves were incubated in 50% TCA (1 mg tissue: 1 μL solution) for 5 minutes then homogenized. Supernatant absorbance was measured at 620 nm, and Evan’s blue concentration was interpolated from a standard curve.

2.5. Data analysis

All data were analyzed by an experimenter blinded to genotype. In the interest of data transparency, individual data points were presented in addition to group mean ± SEM. Two group data sets were analyzed through unpaired t test. Axon diameter distributions were compared through χ² analysis followed by corrected Fisher’s exact tests. G ratios were analyzed through linear regression. Data were analyzed using GraphPad Prism 6; results were considered statistically significant when P < 0.05.

3. Results

Gross changes in SCD-myelinated nerve histology were assessed using light microscopy (Fig. 1A). Berk SS nerves contained a greater proportion of the smallest diameter myelinated fibers (0.5–1.0 μm) and a smaller proportion of 1.5 to 2.0 μm diameter fibers than control nerves; no differences were noted in large diameter fiber abundance (Fig. 1B). Increased numbers of pathological myelin patterns were observed in SCD nerves (Figure 1A, C, D). Myelin sheath infolding and ring separation were the most commonly observed patterns. Thinner myelin sheaths (indicated by increased G ratios of axon diameter/myelinated fiber diameter) were observed in SCD nerves (Fig. 1E). Western blotting revealed increased MBP expression in SCD nerves (Fig. 1F), suggesting that resident Schwann cells are actively myelinating peripheral fibers in SCD so as to replace unmyelinating myelin sheaths.

Gross changes in SCD C fiber histology were assessed using TEM (Fig. 2A). In SCD nerves, fewer C fibers were packaged per Remak bundle (Fig. 2B). Nonmyelinating Schwann cells also occasionally failed to completely ensheathe C fiber axons in SCD nerves (Fig. 2C). Both the nonmyelinated and myelinated peripheral fiber pathology in SCD tissue may have resulted from the increased permeability of the blood–nerve barrier observed through Evan’s blue plasma extravasation (Fig. 2D).

4. Discussion

Many patients with SCD describe their pain symptoms using neuropathic qualifiers. Similarly, transgenic mouse models of SCD display neuropathic pain-like behaviors that are similar to those observed in patients with SCD. Here, we present the first histological evidence that characterize Berkeley SCD mice as a naturally occurring model of polyneuropathy. To date, nerve biopsies from patients with SCD have only been presented in 1 case report; qualitative analysis revealed thinner than normal myelin sheaths, segmental demyelination, and myelin ovoids in the sural nerves of patients with SCD, suggesting a similar widespread myelin instability phenotype in both patients and mice with SCD. However, individuals examined in the case report were also receiving sodium cyanate therapy, a drug with known myelinotoxic effects, and therefore, generalizations about the prevalence of this phenotype in the broader patient population should not be made. Notably, while not apparent in SCD mice, decreased conduction velocity, a possible electrophysiological outcome of reduced myelin thickness, has been reported in SCD patient nerves.

In addition to decreased thickness, the myelin sheath of many SCD mouse fibers appeared unstable; myelin sheath infolding and ring separation, patterns commonly observed in nerve biopsies from patients and mouse models with polyneuropathy, were frequently observed in SCD tissue. Increased MBP expression was noted in SCD nerves suggesting ongoing myelination; validation of these processes should be addressed with additional myelin-associated protein expression profiling. Regardless, when considered with the peripheral nerve sprouting and irregular epidermal innervation qualitatively observed in these mice by other groups, these data strongly suggest that the chronic pain-like behaviors exhibited by SCD mice are generated, in part, through peripheral neuropathic mechanisms. These data are further supported by electrophysiological recordings performed in SCD mice. Functionally identified Aβ, Aδ, and C fibers from SCD mice exhibit decreased stimulation thresholds and increased firing upon receptive field stimulation. C fibers from SCD mice also exhibit high levels of spontaneous activity. Here, we report that in SCD tissue, nonmyelinating Schwann cells sometimes fail to completely ensheathe C fibers contained within a given Remak bundle. This absence of complete fiber insulation may lead to cross-excitability or ephaptic firing between axons, whereby resulting in the increased C-fiber “spontaneous” activity and increased pain-like behaviors observed in SCD mice.
Figure 1. Myelinated fiber pathology in SCD mice. (A) Pathological fibers (Δfiber with myelin sheath infolding; *fiber with unraveling myelin sheath) are observed in cross-sections of SCD sciatic nerve (scale bar = 2 μm). (B) Distribution of myelinated axon diameters in control and SCD mice (χ² P < 0.0001; Fisher’s exact test *P < 0.05, **P < 0.01, ****P < 0.0001; n = 4 biological replicates; n = 100 fibers/biological replicate). (C) Representative TEM images of pathological fibers (*unraveling myelin sheaths; Δmyelin sheath infolding; scale bar = 2 μm). (D) Increased prevalence of pathological fibers is observed in sciatic nerves from SCD mice (unpaired t test ****P < 0.0001). (E) Increased G ratios (ie, thinner myelin sheaths) are observed in SCD fibers of all diameters (linear regression ****P < 0.0001). (F) When normalized to β-tubulin, Western blots reveal increased MBP level in SCD sciatic nerves (unpaired t test *P < 0.05; n = 7–8). MBP, myelin basic protein; TEM, transmission electron microscopy; SCD, sickle cell disease.
In patients with SCD, additional evidence for myelopathy comes from hyperintensities of cerebral white matter observed through magnetic resonance imaging. These imaging abnormalities, which are correlates of demyelination and axonal degradation, may arise from silent cerebral infarcts that are common in SCD, or be subclinical indicators of widespread neuronal instability. Neuronal destabilization in this disease may result from autoimmune mechanisms such as those observed in multiple sclerosis, another chronic proinflammatory disease, or from circulating factors leaking through the highly permeable blood–nervous system barrier demonstrated here in SCD mice. Such circulating factors include lysophosphatidylcholine, a membrane lipid and known demyelinating agent that is increased in serum from patients and mice with SCD, and free hemoglobin, a chemical released from sickled red blood cells that has neurotoxic effects. Although histological examination of patient tissue is needed to validate these preclinical findings, data like these will aid in the development of novel therapeutics that target the root cause of many SCD phenotypes, rather than simply alleviating disease symptoms such as pain.

Disclosures
The authors have no conflict of interest to declare.

This study was supported by research funding from the National Institutes of Health to K.E. Sadler (grant F32NS106789) and C.L. Stucky (grants NS040538 and NS070711).

Acknowledgements
Authorship contribution: K.E. Sadler designed all experiments, performed light microscopy and Western blots, analyzed all data, and wrote and edited the manuscript. T.R. Lewis performed TEM and edited the manuscript. T.B. Waltz performed Evan’s blue experiments, assisted in image analysis, and edited the manuscript. J.C. Besharse performed transmission electron microscopy and assisted in experimental design and manuscript

Figure 2. Unmyelinated fiber pathology in SCD mice. (A) Unmyelinated C-fiber pathology was examined through TEM in cross-sections of control and SCD sciatic nerves (scale bars = 1 μm). (B) Fewer C fibers were contained within the Remak bundles of SCD nerves (unpaired t test *P < 0.05; n = 4 biological replicates). (C) Representative image of SCD Remak bundle in which nonmyelinating Schwann cell has failed to complete ensheath C fibers (potential points of direct contact between fibers; scale bar = 500 nm). (D) Increased permeability of the blood–nerve barrier is observed in SCD mice as measured by Evan’s blue plasma extravasation (unpaired t test *P < 0.05; n = 6). SCD, sickle cell disease; TEM, transmission electron microscopy.
Jessen KR, Mirsky R. Schwann cells and their precursors emerge as
Hillery CA, Kerstein PC, Vilceanu D, Barabas ME, Retherford D, Brandow
van der Land V, Hijmans CT, de Ruiter M, Mutsaerts HJMM, Cnoss
Kohli DR, Li Y, Khasabov SG, Gupta P, Kehl LJ, Ericson ME, Nguyen J,
He Y, Wilkie DJ, Nazari J, Wang R, Messing RO, DeSimone J, Molokie RE,
Garrison SR, Kramer AA, Gerges NZ, Hillery CA, Stucky CL. Sickle cell
Devor M, Wall PD. Cross-excitation in dorsal root ganglia of nerve-injured
Cermenati G, Abbiati F, Cermenati S, Brioschi E, Volonterio A, Cavaletti
Brandow AM, Farley RA, Panepinto JA. Neuropathic pain in patients with
Ballas SK, Lieff S, Benjamin LJ, Dampier CD, Heeney MM, Hoppe C,
Ahmed S, Ali J, Mahdy A, Gadallah N, Hefnawy H. Trigeminal nerve
References
Accepted 19 May 2019

[1] Ahmed S, Ali J, Mahdy A, Gadallah N, Hefnawy H. Trigeminal nerve
electrophysiological assessment in sickle cell anemia: correlation with
disease severity and radiological findings. Egypt Rheumatol Rehabil
2015;42:73.
[2] Balas SK, Leff S, Benjamin LJ, Dampier CD, Heeney MM, Hoppe C,
Johnson CS, Rogers ZR, Smith-Whitley K, Wang WC, Telen MJ;
Investigators, Comprehensive Sickle Cell Centers. Definitions of the
phenotypic manifestations of sickle cell disease. Am J Hematol 2010;85:
6–13.
[3] Brandow AM, Farley RA, Panepinto JA. Neuropathic pain in patients with
sickle cell disease. Pediatr Blood Cancer 2014;61:512–7.
[4] Cermenati G, Abbiati F, Cermenati S, Brioschi E, Volonterio A, Cavalletti
G, Saez E, De Fabiani E, Crestani M, Garcia-Segura LM, Melcangi RC,
Caruso D, Mitro N. Diabetes-induced myelin abnormalities are associated
with an altered lipid pattern: protective effects of LXR activation. J Lipid
Res 2012;53:300–10.
[5] Conran N, Belcher JD. Inflammation in sickle cell disease. Clin Hemorheol
Microcirc 2018;68:263–99.
[6] Devor M, Wall PD. Cross-excitation in dorsal root ganglia of nerve-injured
and intact rats. J Neurophysiol 1990;64:1733–46.
[7] Garrison SR, Kramer AA, Gerges NZ, Hillery CA, Stucky CL. Sickle cell
e mice exhibit mechanical allodynia and enhanced responsiveness in light
touch cutaneous mechanoreceptors. Mol Pain 2012;8:62.
[8] He Y, Wilkie DJ, Nazari J, Wang R, Messing RO, DeSimone J, Molokie RE,
Wang ZJ. PKCβ-targeted intervention relieves chronic pain in a murine
sickle cell disease model. J Clin Invest 2016;126:3063–7.
[9] Hillery CA, Kerstein PC, Vilceanu D, Barabas ME, Retherford D, Brandow
AM, Wandersee NJ, Stucky CL. Transient receptor potential vanilloid 1
sickle cell disease model. J Clin Invest 2016;126:3063–7.
[10] Jessen KR, Mirskey R. Schwann cells and their precursors emerge as
major regulators of nerve development. Trends Neurosci 1999;22:
402–10.
[11] Kohli DR, Li Y, Khasabov SG, Gupta P, Kehl LJ, Ericson ME, Nguyen J,
Gupta V, Hebbel RP, Simone DA, Gupta K. Pain-related behaviors and
neurochemical alterations in mice expressing sickle hemoglobin:
modulation by cannabinoids. Blood 2010;116:456–65.
[12] van der Land V, Hjirans OT, de Ruiter M, Mutsaerts HJM, Crosseen
MH, Engelen M, Majoe CBLM, Nederveen AJ, Groothuis MA,
Fijnvandraat K. Volume of white matter hyperintensities is an
independent predictor of intelligence quotient and processing speed in
children with sickle cell disease. Br J Haematol 2015;168:553–6.
[13] Lee SM, Sha D, Mohammed AA, Assress S, Glass JD, Chin LS, Li L. Motor
and sensory neuropathy due to myelin infolding and paraclinical damage in
a transgenic mouse model of Charcot-Marie-Tooth disease type 1C.
Hum Mol Genet 2013;22:1755–70.
[14] Lewis TR, Kundinger SR, Pavlovich AL, Bostrom JR, Link BA, Besharse
JC. Cos2/Kif7 and Osm-3/Kif17 regulate onset of outer segment
development in zebrafish photoreceptors through distinct mechanisms.
Dev Biol 2017;425:176–90.
[15] Lisney SJ, Devor M. Afterdischarge and interactions among fibers in
damaged peripheral nerve in the rat. Brain Res 1987;415:122–36.
[16] Mayo L, Quintana FJ, Weiner HL. The innate immune system in
demyelinating disease. Immund Rev 2012;248:170–87.
[17] Okuyucu EE, Turhanoglu A, Duman T, Kay, H., Melek IM, Yilmazer S.
Peripheral nervous system involvement in patients with sickle cell disease.
Eur J Neurool 2009;16:814–8.
[18] Orta S, Henry K, Mantuano E, Yamauchi K, De Corato A, Ishikawa T,
Felti ML, Wbrabet L, Gaultier A, Pollack M, Ellman M, Takahashi K,
Gonias SL, Campana WM. Schwann cell LRPI regulates remak bundle
ultrastructure and axonal interactions to prevent neuropathic pain.
J Neurosci 2013;33:5590–602.
[19] Plazty BR, Clum MJ, Marc E, Witkowska HE, Stevens ME, Mohandas N,
Rubin EM. Transgenic knockout mice with exclusively human sickle
hemoglobin and sickle cell disease. Science 1997;278:876–8.
[20] Peterson CM, Tsaris P, Ohnishi A, Lu YS, Grady R, Cerami A, Dyck PJ.
Sodium cyanate induced polyneuropathy in patients with sickle-cell
disease. Ann Intern Med 1974;81:152.
[21] Rigaud M, Gemes G, Barabas ME, Chernoff DI, Abram SE, Stucky CL,
Hogan GH. Species and strain differences in rodent sciatic nerve anatomy:
implications for studies of neuropathic pain. PAIN 2008;136:188–201.
[22] Sadler KE, Zappia KJ, O’Hara CL, Langer SN, Weyer AD, Hillery CA,
Stucky CL. Chemokine (c-c motif) receptor 2 mediates mechanical and
cold hyperalgesia in sickle cell disease mice. PAIN 2018;159:652–63.
[23] Uhelski ML, Gupta K, Simone DA. Sensitization of C-fiber nociceptors in
mice with sickle cell disease is decreased by local inhibition of
anandamide hydrolysis. PAIN 2017;158:1711–22.
[24] Vanderveldt GM, Regan RF. The neurotoxic effect of sickle cell
hemoglobin. Free Radic Res 2004;38:431–7.
[25] Wallace VCG, Cottrell DF, Brophy PJ, Fleetwood-Walker SM. Focal
lyssolecithin-induced demyelination of peripheral afferents results in
neuropathic pain behavior that is attenuated by cannabinoids. J
Neurosci 2013;33:2221–33.
[26] Wu H, Bogdanov M, Zhang Y, Sun K, Zhao S, Song A, Luo R, Parchim
NF, Liu H, Huang A, Adebiyi MG, Jin J, Alexander DC, Milburn MV,
Idowu M, Juneja HS, Kellems RE, Dowhan W, Xia Y. Hypoxia-mediated
anandamide hydrolysis. PAIN 2017;158:1711–22.
[27] Yaghshes S, Matsuaga M. Ultrastructural pathology of peripheral nerves in
patients with diabetic neuropathy. 1979. Available at: https://www.
jstage.jst.go.jp/article/tjem1920/129/4/129_4_357/_pdf. Accessed
February 11, 2019.
[28] Yusuf HR, Atrash HK, Grosse SD, Parker CS, Grant AM. Emergency
department visits made by patients with sickle cell disease: a descriptive
study, 1999-2007. Am J Prev Med 2010;38:S536–41.