LOOSE CONNECTIONS

BRINGING TOGETHER THE VOICES OF OUR COMMUNITY

NOV 2020 ISSUE
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**AUGUST 2020 “COMFORT”**

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She’s prepped and ready, we’re about to start
Okay sweetheart, here comes the easy part
I need you to keep breathing real deep, nice and slow
Your eyes are going to get heavy and it will be over before you even know
You shouldn’t feel a thing, maybe a little burning in your I.V.
There’s just one last thing that I need you to do for me
Count backwards from ten….

TEN.
I said I wouldn’t do this again.
NINE.
Lay on this operating table another time.
EIGHT.
Unfortunately this surgery wasn’t up for debate.
SEVEN.
I hope that things are actually pain-free in heaven.
SIX.
Please let this be a permanent fix.
FIVE.
I’ve come so far, I was dying, now I thrive.
FOUR.
I feel like I’m constantly grieving the life I had before.
THREE.
Trying to accept this new version of me.
TWO.
Being held together by tape and glue.
ONE.
Wake me up when we’re done.
SAFE • VERONICA LEIGH CARR

18x18 inches
Acrylic and pencils on paper

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I created this with people in mind who have experienced recurring trauma over their lives. My intent is to express what that feels like inside, so they know they’re not alone. For people who haven’t suffered profound injury, trauma, or abuse, my intention is to help them better understand what these people feel like. There is innocence and vulnerability buried within even the most hardened, prickly personality.

**The first layer of protection:** The subject, a young adult female, rests in the center of her memories and past, both trapped within and protected by concentric barriers built over a lifetime. Each one is aged on its outside, showing the passage of time from one event to the next. The spots of damage represent trauma, injury, abuse or otherwise damaging events that necessitate the building of another barrier – which is constructed out of whatever materials are available to the young subject. The warm red center is womblike, signifying both comfort and incomplete development.

**Second layer:** A baby blanket, with the first signs of trauma early in her life. As a protective layer it is comforting but quickly damaged.

**Third and fourth layers:** The young subject tries to discourage further pain
by constructing barriers with the only materials she has. First, paper signs with a warning, then a wall of blocks (red, yellow, blue, the primary colors of childhood). When she is hurt again, she is still too young to know how to heal from the trauma, and the barriers remain damaged, the pain still raw.

**Fifth layer:** She is now more experienced at protecting herself. But the wood layer is not enough.

**Sixth:** This time she manages a crude repair in the previous layer, though it remains obviously damaged. Then she builds another outer layer, of concrete, which almost nothing can penetrate. Still she suffers more hurt, but the hardness of this wall protects her psyche better.

**Seventh:** Tired of years of pain, she builds a layer of spikes — becoming not just withdrawn and guarded but sharp, harsh, repelling others before they get to her.

**Eighth:** A row of spikes are effective at preventing harm, but the subject is now unapproachable and lonely. The last layer slowly builds on the outside, rich earth and then lush greenery, forming a more pleasant exterior — as long as no one digs too deep. This is the layer that is presented to the outside world.

I think that deep down, this person knows that she holds the key to her own liberation. But first, she must uncover and process each hurt, allowing it to heal and each barrier to fall away. Then she will feel the warmth and freedom of the world outside — giving herself the gift of a sometimes painful but much fuller existence. ■■■
Bravery

JOSHUA RANSOM

On the darkest hour of the longest day,
a bright light looms over a deafening quiet bay.

Over yonder on top of cold water,
lay a sailboat of which time forgot her.

Drifting along, without a song,
nothing to bore a purpose to ignore
existing to exist nevermore.
Our local heath is stunning in the late autumn sun, taking a slow stroll around with my six year old after school with my camera. Kicking leaves, feeding the ducks, and watching the sun slowly set.
Comfort and Solace

CANDACE WILLHITE

WHEN I THINK OF COMFORT AND solace, I’m sure it’s not what other people would typically expect. There are days when I am in bed; days where I force myself out of bed; days when I’m up and going about my day until time for bed.

From my experience, for myself and my many rare conditions, consistent bed rest is not good, nor is consistent running about. My comfort is being on my heating pad, swallowed in blankets, crime show on, and something to drink on nightstand (water mostly).

Comfort leads to solace but unfortunately at times agony – as you see moving around helps, moving too much hurts, moving not at all does both. Varies day to day. Things I used to do just four years ago are highly unlikely now.

The family I have created understands my conditions. My born-into family is not as aware.

Things I loved doing and once enjoyed, shower me with disbelief, depression, and a broken feeling. What have I become? Who am I? Psychological mind is forever altered as well as physical. I am not the same person now as I was when born. Now I have dos and don’ts I never would have expected. Ever!

Being around others who understand make me feel not as alone in the battle. Making people aware and understand helps too.

I had to do my own medical research in the beginning to find most of my conditions. Unfortunately sometimes you have to when you’re not the “typical age” for things.

If my illnesses were visible, I picture a woman whose whole body was attacked by a tiger, shredded by glass, rods, screws, and tiny daggers. To the world my illnesses are invisible; however, I am not! I want to raise awareness for the “invisible” diseases, because not all are black and white. I have stunned quite a few doctors with my health.

I hope this can reach at least one person who now knows they're not alone.
Exploring the coexistence between pain and beauty. I made this piece as a way to evaluate how I feel as a woman with EDS and combining the beauty of the female body with the fragility I experience represented by the purple and yellow hues.
Invisible is how I would describe myself.
A beautiful commodity to stash on a shelf.

Maybe I am just a different breed.
My body screams from inside and I can feel my heart bleed.

My joints are fighting for their rights while my insides are trying to heal.
I’m trying to make sense of this, I can not believe it is real.

“I don’t understand, what do you mean, I’ve had this all my life?”
It felt like I was in a foreign land, being stabbed with a serrated knife.

My pain is real, it is not in my head, and is rapidly taking its toll.
Rising and smiling to see my beautiful family has become my one and only life goal.

The mere thought of what I’m in for scares the living hell out of me.
Baby steps and one day at a time is how it really must be.

Doctors claim, “There is no cure. You are ill and this is real.”
“I am an active woman, wife and mother, I have no choice but to heal!”

Every joint is loose and one wrong move could set me back for weeks.
The truth is, though, my pain threshold has hit its highest peak.

I can’t explain the daily pain I’m in or what my insides endure.
I’m living in my own torture chamber and they claim there is no cure!

They say, “Be brave, be positive, you don’t look sick, and stop worrying about tomorrow!”
“My God,” I pray, “Please save me from today’s excruciating pain, it is stealing my youth and bringing my life such sorrow. Please God, continue to bless my family, my friends, my strength, my hope, and my faith that I may fight through it all over again – tomorrow.”
There was a little knock at my bedroom door that woke me up. I sat up, winced at the pain, ignored the dizziness, and tried to figure out what time of afternoon, or at this point, night it was. I was slightly panicked that I had overslept for school, but I soon realized someone was waking me up for dinner. I forced myself to stand. My legs were weak. I could feel my heart racing. My back ached. My vision blurred. A pain went shooting through my head. The four-hour nap was nowhere near enough.

I lost my balance a little bit while walking down the steps. I sighed and sat down at the table, body hunched, face sunken. I noticed that half of the family already finished their dinner and rushed around to get ready for their baseball practices.

“I tried waking you up earlier because I wanted you to help me with my homework,” my little sister said sadly, “but mom yelled at me because you were sleeping. I don’t understand why you have to sleep so much. It’s not fair because we are stuck doing all of the chores.”

“I’m sorry, I didn’t feel good,” I said as I tried to play off the tears that filled my eyes.

After dinner I sat on the couch and looked over my assignment book to see what I had for homework. I skimmed over the notes, tests coming up, assignments due tomorrow, a project due next week. I guess I’ll start with the math. After about fifteen problems I felt a migraine coming on, so I packed everything up and got ready for bed. Pain. Dizziness.

The next day in class, I could not focus on the lecture because I was too anxious about getting up and using the bathroom. I hated drawing attention to myself and I had already gone to the bathroom once this class period. I dreaded getting up again in the middle of class and having the teacher sign my pass. Finally, I could not hold it any longer and I stood up to leave. Pain. Dizziness.
I walked back in and realized that everyone was packing up their things. The class was walking up to the computer lab to work with Excel.

Great.

I gathered my stuff and walked the opposite way down the hallway.

“Hi, I need the elevator again, please,” I asked the office ladies.

“You’re gonna have to hold on. Didn’t we just get the elevator for you last period anyway?” she said.

“Yeah, I’m so sorry my class is — is — is walking up to the — the — um, thing you know — the computer lab.” I struggled to find the words. Ugh why did it come out like that I sound like an idiot.

The woman rolled her eyes. If only they knew how much I did not want to take the elevator and how much I did not want to bother them.

The janitor walked with me to the elevator and when we got in, I laid my head against the wall and closed my eyes. Ugh four more hours I can do it. Four more hours until I can lie down.

When I finally got to the classroom, I asked the kid next to me what we were doing because the seats by my friends were already taken. He explained we were doing some problems in Excel from our textbook.

I stood up. Pain. Dizziness. I walked over to the window where my extra book is kept since I could not carry my own. Once again, it was missing. A migraine started forming.

“Sorry to bother you, but my extra book is not where it is supposed to be again,” I said to the teacher.

“Oh yeah, I had to give that out to one of the new students, sorry I forgot to replace it. Can you just share with someone?” he asked.

“Sure, I guess,” I murmured anxiously. My classmates probably think I’m so annoying.

“Also, while you’re up here, when did you want to make up that test you missed? I wanted to give it back to the students because I already graded them,” he said.

Great now all of my classmates know I’m the reason they can’t see their grade yet.

“Oh, probably Friday, because I am making up my math test — ummmmmmm — tomorrow and meeting with someone on — uhhh — Thursday I think, about what I missed in class,” I tried to spit out while another wave of brain fog started to consume my head making it difficult to think and speak.

“Oh, wow, that’s a lot of make-up stuff. It seems more complicated than just doing it when it is due, huh,” he said.
Wow. WOW. Never thought of that.

I was not even surprised by the comment, I just felt defeated once again. I wish it did not have to take being directly affected by something for people to have empathy.

Instead of going to my next class, I found myself drudging to the nurse’s office, tripping over my feet and bumping into the walls like I was drunk. I squinted at the light when I walked in. My headache was getting worse and now I felt nauseous.

“Hi hun, the second room is open. Do you feel like you need to go home?” she asked.

I nodded my head and walked over to her fridge. I grabbed a pack of saltines and flopped down on the bed. Ugh that was too fast. Pain. Dizziness.

I was confused at why I started to cry. But many thoughts started flooding in while I was lying there, staring at the ceiling.

Why am I so angry? I feel like I have a right to be angry because I’m not living a normal life. This is unfair. Why don’t I have it as easy as my peers. It’s so hard for me to do normal things. I can't even walk up the steps without needing to sit down and take a break. That’s pathetic. I’m pathetic. The people at this school are pathetic. Why is this happening to me. Why don’t people even feel bad for me? I’m sure my teachers would take my accommodations seriously if I was in a wheelchair, if I was disfigured, if I had cancer! Why am I thinking like this? I should be so grateful that my condition is not worse than it is. But I wish I could just shake these people and scream that I’m not making it up! I’m not taking advantage of the system! I’m in pain! It’s extremely hard for me to live my life everyday! Where is my recognition? Stop thinking like this! I’m a horrible person. I’m so much better off than others that have more of a struggle. Calm down. CALM DOWN. I don’t need pity to feel like my struggles are validated.

“Your mom’s here,” the nurse whispered into the room.

Pain. Dizziness.

Thank God, get me out of here.
“Why are your eyes so red?”
My eyes are red because I am in agony,
But you’d never be able to tell
Because although I deal with it everyday,
You’d never care enough to notice.

“Why are your eyes so red?”
My eyes are red because I am lonely,
But my lonely does not mean I’m alone.
My lonely is forcing a smile in a grocery store,
Because my anxiety tells me that
no one likes someone unfriendly.

“Why are your eyes so red?”
My eyes are red because I love you,
But my love is hidden and alone.
My love is unrequited and rightfully so;
Who could ever love me
If I can’t even love myself.

“Why are your eyes so red?”
My eyes are red because I am scared,
But I’ll never tell you that.
Scared because I feel hopeless,
And I overheard my neighbors saying that
people who are hopeless have no future.

“Why are your eyes so red?”
My eyes are red because they leak,
And the saltwater they leak is painful.
Painful because I know that I need help,
But I was taught to be independent
So asking for help is too hard.
Image description: Greys, whites, blacks, greens, and a small red dot color this large off-white cardstock painting oriented in portrait. There are grey and green vertical smears made with the feet and some identifiable footprints, toe marks, and a circle where the ball of the foot spun. There are also two splattered perforated lines of paint going roughly down the center vertically, one each in black and white.

36x24 inches
Acrylic paint, charcoal powder, and ink
December 2019
Mikhail Bakunin said, “The urge to destroy is also a creative urge!”

As an artist, I process the world through making and being able to use my body to do so. In 2019, I found myself in an acute state of impairment and pain: lumbar and cervical herniations, nerve impingement, radiating bilateral leg and arm pain, thoracic outlet syndrome, limp wrists, headaches, and ankle and knee pain. My usual ways of creating were untenable and without a way to express and discover meaning, my quality of life was at one of its lowest. I could hardly use my hands, cycled between icing either lying down or pacing, and required extensive assistance to make it through the day.

I attempted new ways of creating: audio transcribing prose, drawing while lying down in a rescue position for my spine, drawing with wrist braces on lightweight paper while walking around town, molding small lumps of clay while pacing.

Offering a helping hand, a painting mentor helped gather materials and taped large papers to the floor so I could paint with my feet and use kinesthetic movements as a guide to mark making. I remember feeling torn between the desperation to express myself and the cries from my body to focus on form — proper foot placement, soft knees, core drawn in, ribs anchored, neck in neutral — feeling plagued by the familiar conflicts and trade-offs between physical self-maintenance and life experience and opportunity.

Learning to navigate this conflict by honoring and evaluating the benefits and consequences of both decisions is a skill I am working on in order to manage the physical and psychological effects of EDS. This also means learning to both repair the damage of the sacrifices and celebrate the self-care and nourishment in however I choose to engage in or refrain from opportunity. Having EDS is to share a body and a spirit that do not harmonize and to live in the gap in between, trying to make this liminal space a forever home.

It has been a year since I made Misaligned Spines and I have progressed since thanks to regular physical therapy, body work, holistic treatments, medication, my blooming meditation practice, fierce friendships, psychotherapy, the hours I spend everyday managing my (re)injuries, and a whole lot of patience. This was never the life I had imagined or hoped, but it’s somewhere caught in between that and the limitations of a body I am learning to work with. A difficult but integral in-progress lesson for me is that my body is not trying to destroy me; the breaking days and painful movements, though dysfunctional, are still sincere attempts at motion and supporting my curious spirit. My body is my friend; it is like me, trying to support itself with the resources it has.

For more on her work, visit www.nweinart.com
Image description: Free colorful squiggles made with pencil dance in a background of mostly blues, salmon, and rich yellow. Vibrant colors — all shades of blue, lime green, rich yellow, bright red — smoothly smear down the page vertically. Circles of color can be seen at the top of the smear where the paint was applied to the page. The painting evokes something of release, freedom, sensation, and movement.
“What is escape”

J. FISHER POETRY

What is escape
What does that even mean
Because the escape that I want
Is to escape being me
Inside this body
Is a living hell
But I wasn’t given an option
For anything else
But days like today
I want to crawl out of my skin
When the pain is this bad
I want to tear off my limbs
My ears are ringing
Like an incessant alarm
Except I can’t turn it off
When the noise causes harm
Oops, there goes my hip
Better pop that back in
Then I’ll have to do it again
And again and again
But I keep moving forward
With a smile on my face
Because I have no other choice
I don’t get to escape
Some days are hard
And some days are harder
But the only choice I have
Is to be a survivor
BREADCRUMBS ARE THE SMALL positives that come out of the chronic illness battle. For me, it is so important to try and find these breadcrumbs, however tiny, in every terrible moment. It’s how I deal, how I stay afloat in what feels for the most part like a pool of misery.

Right now, my breadcrumb is being grateful for the person that my chronic illness has moulded me into. Without this battle I would never have become the version of me that I am today. Something tells me there are many others out there who are proud of what their struggle has made them, and if you’re not, perhaps you should give yourself some credit.

You see, when you are a zebra, you are forced to examine yourself internally on a level that most would never comprehend or experience: being constantly asked to justify your pain, explain your situation to those without comprehension or empathy, being ignored, being shamed and being told countless times it all in your head. Internally, I assure you, that doesn’t go unexamined. If you are like me, you spent an inordinate amount of time in self-reflection mode, taking a deep dive into your own psyche, wondering if they are right. Analyzing every pain, every symptom, every moment of suffering and grabbing yourself by the metaphorical collar with a shake, demanding “why are you being like this, what is really going on here?” If you are like me, you did this over and over for years and you now know, without a shred of doubt, that none of this is in your head.

You learn so much more on this journey of self-exploration and this is something we should celebrate. I believe it leaves a permanent mark on you, it becomes part of your soul and changes your personality in so many positive ways. That sort of deep internal reflection brings to the surface something special, almost like a sixth sense, a superpower of sorts perhaps! Sure, there are the not-so-good outcomes too, but the good far outweigh the bad and we have enough bad in our lives already, so let’s focus on the good.
The good outcomes, the positives that emerge, are abundant and powerful – you will become strong beyond belief, both mentally and physically. You will be able to handle criticism and cruelty like drops in the ocean, water off a duck’s back as they say. You are able to stand up for yourself when others wouldn’t.

You learn to speak up when you are being mistreated. You ask questions without fear of judgement. You push back when you are pushed. You learn not to underestimate yourself. You learn to take on every day, every battle, head on and without fear. You learn to use these powers for good, and now you help others who are struggling to find their voice. You learn to be gentle with others who could be suffering on the inside just as you are.

You learn to scream in silence. You learn to cry without tears. You learn to smile when inside you feel numb. You learn to pull yourself together, when your body falls apart. You learn to smile and nod as people pick apart your condition without any actual concept of what they are saying. You learn that people’s opinions are mostly not worth your time, they are certainly not worthy of causing you more pain. You adapt in evolutionary ways to accommodate the broken body that nobody sees, so you can pretend to be just like everybody else, even if just for a little while.

You learn to read people like a psychic. You understand every look, every twitch, every tone. You know when someone is worth your time and when they are not. You know when they are open to using their skills in helping you through your journey or when they are angered by the unknown you present them with. You know that there are far more disingenuous people out there than everyone else realizes. You know that the world is not as nice of a place as it should be, but you know you are strong enough to find your place in it. And if there isn’t one, you are damn sure you are strong enough to create one that's right for you.

You learn to stand tall, even when you feel small. You learn that every failure is still a step towards growth and you are strong enough to fail over and over again if you have to. You learn that there are no easy answers, no predetermined outcomes, no miracles, and no favors. But that’s OK, nothing worth having ever comes easy and you know your abilities, you know what you can accomplish despite all your barriers.

We've learned to fight battles that shouldn't exist. We've cried so many tears there are rarely any left. When we allow ourselves to break, it's only for a few private minutes. We've learned to collect ourselves, reattach that forced smile and light-hearted confident persona. We've learned that we are just as worthy and just as valuable as everybody else, even if other people don’t see it yet. We've
learned it is OK to ask for help, it doesn't diminish our abilities or mean we're giving up. We've learned to breathe in a whole new way: breathe to fight pain, breathe to calm ourselves, breathe to allow us to dig deeper within ourselves to come out swinging when we need to. We've learned to laugh: at ourselves, at the sheer ridiculousness of our situation and the roller coaster ride that is our every day. We often feel like life's practical joke, but we've learned to laugh with and in the face of it.

Every day we get through builds our strength and our determination. If you look closely, really closely, you might catch a fleeting glance of who we really are. You might see the storm that's rolling through our bodies, silently just under the surface.

We're suffering in every way imaginable, but we're warriors. We're fighting a battle you can't see. And the fact that we're standing before you now means we still, in at least some small ways, are winning.
“I’m sick”

MEGAN MCCARTY

I'm sick
Of the steps that block my path
And lift others higher who were only lucky enough
To be born THAT way.

I'm sick
Of the beeping monitors
That lead me through my youth
instead of buzzers
exclaiming my victory
because my body COULD.

I'm sick
Of the tears I weep
When the burden to bear
somehow ALWAYS is theirs.
I'm sick of the constant change
And the cold of each day
And shoulder of those I
USED TO know.

I’m sick

And I’m SICK of it.
EDS, I Hate You

VERONIQUE LANGLOIS

You are too close for comfort
Always want to be in charge
You pin me against the wall
And never leave me in peace

You are stalking me everywhere I go
You watch me like a hawk
You are a mean monkey on my back
You are the elephant in the room

You always open a can of worms
I am tired of your charley horses
You are my biggest pet peeve
I wish I could just quit you cold turkey

You are a pain in my neck
A chip on my shoulders
A constant pain in my ass
You make me weak in the knees

You hit me like a ton of bricks
You slap me in the face
You keep stabbing me in the back
And you tear me apart

You are a thorn in my side
You make me walk on egg shells
I am always waiting for the other shoe to drop
When all I want is to be in someone else's shoes

Your constant grip made me a twisted sister
You make my head spin
You take my breath away
And I can't get out of your straight jacket

You are the mountain I can't climb
You are the black cloud over my head
You always rain on my parade
I never get to see cloud nine

You are an annoying back seat driver
In my rollercoaster ride
You cost me an arm and a leg
You add insult to my injuries

EDS you suck
WANT TO WRITE ABOUT AN ABDOMINAL aortic aneurysm in a man, whom I will call F. He wanted to find out if he had an aneurysm because several men in his family had died from internal bleeding. He has periodontal Ehlers-Danlos syndrome (pEDS).

F. was 73 years old, and had been referred to a doctor about a lump he had on the left side of the chest. The doctor told him, “That is nothing, it is only fat.”

F. asked, “Why did I pay money to be told it was just fat? Would it be possible for you could check my abdominal aorta?”

“Yes, I can do that,” said the doctor, who examined his abdominal aorta with ultrasound. F. was told, “Oh, it was rather big.”

F. asked, “Can you please write that to the doctor who sent the referral?”

When the doctor who had sent the referral saw the results, she told F., “The question of the lump was answered, it was nothing, so this is finished.”

But F. said, “Please send a referral to the vascular surgeon, since I have pEDS.”

F. received his referral, and he was checked every six months. The abdominal aorta continued growing wider, and when it was 65 mm he was operated on. A second surgery was performed because of a big leakage, and a third surgery repaired the narrow aorta to the left leg.

It was solely his own questions that brought F. to being examined, followed, and eventually successfully treated with surgery.

Since then F. has been fine and has started exercising and goes to the wood picking berries. This is comfort!
Pain!

My pain is real!
And it doesn't show race, color, religion, or culture!
I may look OK and smile!
But actually no one will understand my pain!
Only those suffering from pain will understand!
So I beg thee!
Listen to those living with pain!
As it takes a lot of courage, hope and faith!
To face everyday!
I feed my mind positive thoughts!
To prevent myself going insane!
The Ehlers-Danlos Society is a global community of patients, caregivers, medical professionals, and supporters, dedicated to saving and improving the lives of those affected by the Ehlers-Danlos syndromes, hypermobility spectrum disorders, and related conditions.

We are proudly working to provide global learning conferences, collaborative research and education initiatives, awareness campaigns, advocacy, community-building, and care for the EDS and HSD population.

Our goals are worldwide awareness – and a better quality of life for all who suffer from these conditions. Research is at the center of what we do, so that one day we will have a cure.

OUR STRENGTH BEGINS WITH HOPE.

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