Area families discover innovative treatment programs and a network of support at JMU

HOPE for AUTISM

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PHOTOGRAPHS BY MIKE MIRIELLO ('09M)
HOPE FOR AUTISM

When Laurie (’15M) and Robert Weese retired from the military, they expected to move. But having two sons diagnosed with autism, they considered their options carefully. “We started checking out locations — Tennessee, New Jersey, Indiana, Montana … all over the place.”

While traveling through the Shenandoah Valley one day, they stopped at the Rockingham County School Board offices, where they met Scott Hand (’79, ’80M), director of pupil personnel services. He took them to visit county schools. What the Weeses found was a community supportive of autism and engaged with a university replete with services and opportunities to help their sons.

It’s not uncommon for parents to seek out a school district that will support their child and then move there, says Keri Bethune, coordinator of the autism certificate program at JMU, because support for children and families is essential in addressing autism and its many ramifications.

DIAGNOSIS: MYSTERY
First identified in 1911 as a subgroup of schizophrenia, Autism Spectrum Disorder is a developmental disability causing social, communicative and behavioral difficulties. It became a separate diagnosis in the 1940s, but until the 1960s parents were routinely advised to institutionalize their children. The assumption was that nothing could be done for children locked in their own minds, some without the ability to speak and many without the skills to interact socially.

Autism has been blamed on everything from a lack of mother’s love to genetics to environment to vaccines — the latter, a myth discredited by multiple scientific studies. Real progress in addressing autism began in the 1980s, when treatment options emerged and parental pressure, allied with professionals, began to unlock a diagnosis that had created more mystery and anxiety than clarity.

Gay Finlayson (’76) understood the anxiety. More than three decades ago, she and her family were living with a child they did not understand. The only diagnosis Finlayson received for her daughter, Marit, was Pervasive Developmental Disorder Not Otherwise Specified. In other words, no one really knew what was wrong.

Finlayson, though, had an advocate in her mother, Elizabeth Finlayson, founder of JMU’s Bachelor of General Studies program (now the Bachelor of Individualized Studies). “My mother’s connection to the world of higher education was important,” Finlayson says. In other words, no one really knew what was wrong.

Finlayson’s answer was to advocate for her family, which soon expanded to three children, including a son also diagnosed with autism — a coincidence that is not uncommon. Her personal journey became one that thousands of parents travel in search of help for their children.

THE PROMISE OF SUCCESSFUL TREATMENTS
For many years, insurance companies refused to pay for autism services, offering a single excuse: There is no viable treatment. Parents disagreed. Research in applied behavioral analysis, along with extensive and growing anecdotal evidence, convinced them that treatment was indeed possible.

“What happened,” says Trevor Stokes, professor of graduate psychology and a licensed applied behavioral analyst, “is that the parents who were the consumers of services … could see that it was effective.”

When pediatricians diagnosed autism and prescribed ABA treatments, he says, insurance companies, persuaded by research — much of it emanating from universities — were compelled to comply.

Despite the acknowledgement that ABA worked, there were other limiting factors: Who, for instance, would provide the prescribed treatment, especially as the number of diagnoses skyrocketed? According to the Centers for Disease Control, today 1 in 68 children is identified with ASD, an increase due in part to greater public awareness and an expanded definition. In 2013, the American Psychiatric Association revised the Diagnostic and Statistical Manual of Mental Disorders to lump together several diagnoses, including Asperger’s, a mild form of autism, under the single Autism Spectrum Disorder.
The prevalence of autism has created a need for, if not a shortage of, ABA professionals, Stokes says. He adds that the Affordable Care Act has expanded coverage of mental illnesses in general, plus this year the Virginia General Assembly upped the age limit for required services from 6 to 10 — a change set to begin in fiscal 2016.

To address the need for more ABAs, five years ago JMU developed a collaborative program within the College of Health and Behavioral Studies that prepares graduates to become licensed ABAs. The designation, granted by the commonwealth’s Board of Medicine, requires 1,500 hours of supervised practicum work followed by a board certification exam. Steered by JMU’s three licensed ABA professionals on faculty — Stokes; Keri Bethune, who also serves on the governor’s advisory board on licensing ABAs; and Dan Holt of the psychology department — the program is an integral part of a large network of services and opportunities available to the autism community through JMU.

**A NETWORK OF HELP**

Every week, Garrett Weese, who first came to JMU as a nonverbal 7-year-old, visits Occupational Therapy Clinical Education Services, a clinic within the Institute for Innovation in Health and Human Services. Over the past six years, he has made slow but steady progress. He makes eye contact now, his mother says, and has autism services. These include an ABA clinic under IIHHS and a Speech-Language-Hearing Applied Laboratory, part of the Department of Communication Sciences and Disorders. Each provides a different therapeutic approach.

These different approaches converge in the university’s Interprofessional Autism Clinic. Also under the umbrella of IIHHS, IPAC offers blended autism services to the community. Stokes explains: “IPAC is very innovative because we’ve brought together people from different disciplines — from ABA, from psychology, occupational therapy, and speech and language pathology. We can do the merged treatment at one time … which is not a traditional approach, but this is the creative part of what we do.”

At the same time, IPAC provides a fertile experiential learning environment for JMU students who will eventually work with children on the spectrum. One such opportunity is JMU’s summer autism camp, which is run by occupational therapist Liz Richardson and Marsha Longerbeam, a speech and language pathologist.

While the benefits to children with autism are indisputable, there is added benefit for parents. “When [parents] come in for an appointment, they don’t just drop the child off,” Stokes says. “There’s an expectation they’ll be involved in some way. As a result, parents learn strategies and therapies to help their children that might otherwise take years of trial and error.”

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THE EDUCATION COMPONENT

“No child with autism is like another child with autism,” Bethune says. “It is very much a disability where educational plans, behavior plans and strategies all have to be individualized.”

Bethune, who coordinates the behavior specialist concentration within the College of Education, is a regular presence in local elementary and secondary classrooms, supervising JMU students and teachers who work with children with autism.

“It’s really, really important because we know that autism is one of those disability areas where kids could look a million different ways and be served in a variety of settings,” Bethune says. “They could be in the general education classroom, with no extra support, all the way to a separate school, depending on the level of need of the student. What we know is that getting good evidence-based interventions early makes a very big difference in the outcome for these students.”

A year ago, COE began offering a one-year course sequence leading to an autism certificate — an extension of the university’s traditionally strong exceptional education program.

“The cool thing about the autism certificate program,” says Bethune, “is that it’s not just education students.” It is open to students in all majors and is offered at both the undergraduate and graduate levels. Right now there’s a waiting list to enroll.

The upshot of this kind of collaborative education is that JMU is preparing students in ways that create a kind of autism advocacy throughout the commonwealth.

Scott Hand represents the hundreds of educational professionals coming out of JMU whose impact in local schools is significant. In Rockingham County, autism programs employ the Competent Learned Model, which is based on the principles of ABA, direct instruction and precision teaching.

Robert Weese, Garrett’s father, is especially appreciative of this impact. Before coming to the valley, he would drop by Garrett’s school and find him isolated in a classroom. “Educationally he was being left behind because the teachers were not required to learn. Here in Rockingham County, they are encouraged to learn about autism, and JMU helps that.”

TANGENTIAL OPPORTUNITIES

Beyond IPAC, the campus clinics and school systems full of JMU-educated professionals, other services and opportunities are available to families. Some are associated with Madison directly while others are tangential — like music.

Laurie Weese’s weekly calendar often includes a visit to Studio B Music for Living LLC, a private music therapy studio in Rockingham County. “Garrett loves to sing,” she says.

“Science tells us that engaging in a musical experience activates almost our entire brain,” says Brianna R. Priester, a licensed music therapist, instructor of music at JMU and owner of Studio B. “Individuals who may not be able to verbally communicate may be able to sing. The music works around the language barrier and spreads throughout the brain to activate different triggers.”
therapy can also help with the development of social skills, daily-living activities like chores and self-care, emotional regulation, stress and anxiety management, and self-expression, she says.

Additionally, children with autism benefit from physical and sensory experiences. That is the rationale behind JMU kinesiology professor Tom Moran’s Project Climb, an outreach program of adapted physical education for children with disabilities, including autism.

John “Jack” Martin, a utility locator in JMU Facilities Management, had a similar idea when he, along with his wife Becky (’74) and retired educators Sue (’78) and Philip (’79) Hutchison, developed a baseball program for children with disabilities in nearby Bridgewater, Virginia. Neither official nor clinical, Martin’s program nonetheless provides something extra: “It gives Garrett a chance to feel like an ordinary boy,” Robert Weese says.

**A FAMILY MATTER**

Ask Allison Lindsay (’16) what is wrong with her brother and you’ll get a swift answer: “Nothing. He’s just different.”

Autism has always been a part of Lindsay’s life. Her older brother Sam and younger sister Mikela both have autism. While Mikela falls on the milder end of the spectrum, Sam, who also has epilepsy and some degree of cerebral palsy, is profoundly affected. To Lindsay, though, he’s just her brother.

Lindsay knows firsthand one truism about autism: It impacts the whole family. “I remember going to kindergarten,” she says, “and I had no idea that not everybody had a brother like Sam … but I learned that that wasn’t the case. But I am so grateful to have him. Our family is a very chaotic, fast-beat family, and I think Sam gives us the opportunity to step back. He teaches us patience, and he teaches us just to be empathetic and aware. He’s my favorite person.”

Like so many people who are close to autism, Lindsay sees the disability but also the person. Autism is part-handicap, part-barrier, she says, but it can also come with astonishing talents and potential. One of those is intelligence. “You see a whole range of intellectual levels,” Stokes says, from incidences of autism with intellectual disability “to people on the spectrum whose intelligence is very high.”

Understanding the family dynamic and recognizing a need to support local autism families prompted two JMU professors, Debi Kipps-Vaughan in graduate psychology and Julie Strunk in nursing, to organize a support group for families, especially those with adolescent children on the spectrum. “Teenagers with autism have a unique set of needs,” including independence, sexuality and transition planning for when they leave high school, Kipps-Vaughan says. Now families can come together to support and learn from each other.

Another positive outcome from the support group is the benefit to JMU psychology students, who provide child care. “They get to see and interact with whole families,” Kipps-Vaughan says, “something they could never learn in a classroom.”

**WHAT THE FUTURE HOLDS**

Often overlooked by the media, and to some degree by public opinion, is the fact that, as Finlayson says emphatically, “little children with ASD grow up to be adults with ASD.”

For professionals like Bethune, it’s a concern. “It’s a hot topic,” she says, “because the supports for adults look very different than the supports for children. We still have to have transition services to figure out what’s going to be next for that student. … There’s no mandate for us to continue. So there are a lot of different options. For example, if the child wants to go to college, transition services should help with the application process and then the child has to go through that university’s Office of Disability Services [to] get whatever accommodations they need. They have to advocate for themselves. So self-advocacy is really, really huge.”

Self-advocacy is exactly what JMU professor of computer science Brett Tjaden encourages the dozen or so ASD students he advises every year to practice. The students are referred to him by the Office of Disability Services.

“One of their big challenges,” Tjaden says, “is social interaction. That’s hard enough for most

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By any measure, Micah Hodges ('15) and Ted Aronson ('16) have been successful at JMU. Micah, a gregarious music industry major from Glen Allen, Virginia, performed with the Marching Royal Dukes, joined a social fraternity, and this fall is a FrOG — First Year Orientation Guide — helping JMU freshmen adjust to college life. Ted, a nursing major from Arlington, Virginia, competes on JMU’s archery team, participates in several Jewish social organizations, served as president of the Student Advisory Council for Nursing, and has completed an independent study that his professor calls “revolutionary.”

Both have autism. Diagnosed as a young child, Ted benefitted from early and extensive interventions that included speech and occupational therapies.

“All through elementary, middle and high school,” Ted says, “I was much more focused on academics instead of being friends with people, having peers. When I came to college, and even the last part of my senior year of high school, I thought that it would be more beneficial to shift from being purely academic to having school also be a way of learning social skills.”

Micah’s experience with autism is similar, although his diagnosis came much later. As a high-school student, he knew something was wrong, but he didn’t know what. Once he discovered it was autism, he says, “I was devastated. How do I have autism?” Eventually, though, Micah accepted the diagnosis. “Truthfully,” he says, “I do have to live with Asperger’s for the rest of my life, but the thing is, how you cope with it.”

For Micah, real progress occurred when he came to JMU. “My first challenges were trying to make a group of friends. I found myself very isolated, lonely as a freshman. I would sit alone at lunch every single day. I would always have my head down. It was very difficult to talk to [others]… especially for me, as a minority student as well.” That changed in the spring of Micah’s freshman year. “I had rushed a fraternity called Phi Mu Alpha Sinfonia, which is a men’s social fraternity, and I had gotten a bid. … I became a brother at the end of spring.”
Both Micah and Ted credit JMU’s friendly and open community for helping them carve out successful college experiences despite their disability. Ted chose Madison in part because JMU “embraces being a whole person, like having that social aspect, going to football games, but also being academic,” he says.

Nevertheless, college life can present social challenges for students with autism. “There are still times when I’m talking to people and I’m thinking of what to say, and I don’t know the correct response when someone says something — what’s socially acceptable, what’s not,” Ted says. “There are still small blurred lines in there, but I try to calculate it as best I can … For someone with Asperger’s, we don’t understand the social dynamics of a situation, but if we can understand the people — hence their motives, their view of the world — we can have a better understanding of it, so surrounding yourself with different sorts of people would help accomplish that.”

**Creating Success**

Matt Trybus, assistant director of the Office of Disability Services at JMU, says that Ted and Micah’s experiences are not atypical. During high school and before, he says, the responsibility for accommodating any disability rests with schools, often with significant parental involvement. When a student enrolls in college, however, managing and seeking accommodations for a disability like autism becomes the student’s responsibility. It can create a barrier: How do you get a student who has trouble communicating to do this, including disclosing their disability?

“The system asks people to do something they may not be ready to do,” Trybus says. The solution, “is trying to move the student toward more independence as they’re coming [to college] because they’re going to have to be self-disclose in order to receive accommodations and navigate college.”

Trybus says the students who are most successful have meta-cognitive awareness — they know their strengths and weaknesses, accepting both, and when barriers present themselves, they figure out how to navigate.

“They’ve integrated their disability into their identity,” he says. “That’s not the definitive quality of who they are, but it’s who they are. Those with visible disabilities don’t have any of the integration issues because everyone sees it … You can’t hide it.” Autism, on the other hand, is often an unseen disability.

One pattern Trybus sees is a conflict in students who do not seek help initially. They face a conundrum: Do I suffer the perceived stigma of autism or do I seize the opportunity to re-invent myself? It often comes down to the question of how much a student wants to succeed.

Trybus is aware of about a dozen JMU students who self-identify as having autism. Among the accommodations he can provide these students are additional time and better surroundings to help with test taking. “My advice [to students with autism] is more to know what their strengths are … what they like, what they are passionate about — and where their limitations are,” Trybus says. It’s also helpful, he says, “to know what things, what people and what strategies have been helpful [in the past] … and know what their obstacles are to asking for help.”

“We want all students to succeed at JMU, and that includes students with ASD.”

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**The Gift of Autism**

Until the mysteries of autism are unlocked, families affected by the disorder will continue to focus on the positives.

“It bothers me when people kind of pity me; ‘Oh, I’m so sorry you have an autistic son,’” Laurie Weese says. “But I say, ‘He’s great. He’s just different. He doesn’t talk … but he’s a great kid.’

Siblings of children on the spectrum exhibit a special empathy. “I am not a naturally patient person,” Lindsay admits. “But Sam has taught me that everybody has setbacks, whether it is an experience or a disability, and he makes me so much more aware of that. I just love him so much.” Lindsay has told her parents unequivocally that she intends to take care of her brother for the rest of his life.

The same is true of Garrett Weese’s brother Connor. “Connor is a wonderful kid,” Robert Weese says. “He’s the youngest, but at the same time he has big shoulders. He’s had to help a lot, especially with Garrett. It’s easy for Laurie and me to forget he’s the youngest because we rely on him to help. I thought it would be a problem, but now he’s enthusiastic. If we go roller skating, he says, ‘I’ll go with Garrett,’ and he loves doing it.”

Families like the Weeses and the Finlaysons would agree that there is a certain joy in embracing ASD individuals for who they are and what they can offer. Perhaps that is a lesson autism can teach us all.