Brookes E-Pak Series, #2:

A
utism

featuring material from
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Autism is a developmental disability that typically appears during the first 3 years of life. Classified as a neurological disorder that affects the functioning of the brain, autism affects the normal development of the brain in the areas of social interaction and communication skills. Children and adults with autism have difficulty with communication, social interaction, and leisure or play activities.

**Why Autism as a Spectrum Disorder?**

Major advances have been made since the 1980s in understanding the social and communication difficulties of children with autism. Because individuals exhibit varying levels of impairments, the disorder is classified along a spectrum. The autism spectrum consists of three core features: impairments in social interaction, impairments in verbal and nonverbal communication, and restricted and repetitive patterns of behavior. However, not everyone with autism has all three impairments, and even individuals who do have all three vary drastically in the severity of their impairments.

Here’s a closer look at the three core features of autism:

**Impairment in social interaction.** For children with autism, this is marked by impaired nonverbal behavior, lack of spontaneous sharing, socioemotional response, and/or failure to develop peer relationships.

**Impairment in communication.** This is represented by a delay in or lack of development of spoken language and gestures, the ability to initiate or maintain conversation, repetitive and spontaneous use of language, and/or lack of pretend play.

**Restricted repertoire of activities and interests.** Children with autism are often preoccupied with patterns, inflexible to routines, show repetitive movements, and/or have preoccupation with parts of objects.

The symptoms and characteristics of autism vary by person as well as disorder, from mild to severe, and fall under the umbrella term Pervasive Developmental Disorders (PDD). The following five PDDs make up the autism spectrum:

- **Autistic Disorder:** This is the most common PDD. The word autism was first used to describe a social aloofness and “preservation of sameness” noticed in a number of young children in the 1940s. Many of the features observed then are still observed today.

- **Asperger Syndrome:** Children with Asperger syndrome have the same issues as those with autistic disorder but do not have severe impairments in language development. Children with Asperger syndrome have significantly stronger abilities with rote learning, spelling, and vocabulary.

- **Childhood Disintegrative Disorder (CDD):** With this rare condition, children develop typically until between the ages of 3 and 5. Then, their language, social skills, play, motor skills, and even bowel and bladder control decline. The decline eventually stabilizes, and the child will then have symptoms of autistic disorder.

- **Rett Syndrome:** This primarily affects girls and is caused by mutations of a gene that is essential to early brain development. This is a neurological disorder characterized by loss of purposeful hand use, spasms in the legs, seizures, small head size, and mental retardation.

- **Pervasive Development Disorders-Not Otherwise Specified (PDD-NOS):** This term is used to describe a child whose behavior resembles autistic disorder or Asperger syndrome but may be too mild to truly diagnose. When the child has problems with social learning, however, this diagnosis is often made.
The symptoms in each of these areas can overlap, although the disorders do not. Children and adults can exhibit any combination of behaviors from any area, making it possible for two children with the same diagnosis to exhibit different behaviors, reinforcing why there are no clear distinctions among these five disorders.

**Why early diagnosis?**

Early diagnosis can be the key to a brighter future for a child with autism. The earlier a child is diagnosed, the sooner he or she can benefit from the intervention programs available. Diagnosing a child at an early age not only helps the child develop, but also can be a comfort to the child’s parents. Early diagnosis and involvement with early intervention services gives parents the opportunity to grow with their child and helps them realize that they are not alone. Remember, help is available.
When I look back over my own experiences of searching for help and support in the early days of worrying about Tariq, I see that much of what I went through was fairly typical. At first, I believed that if he could just start talking again then everything would work out from there. When something is wrong with your child, you often feel like a little child yourself, and you want your mom and dad to help you—to rescue you and protect you from the hurt that you are feeling.

My mother and father would counsel me to be patient and tell me that things would work out. My mother would tell me that she was praying. They would ask me how Tariq was doing, as if he had the flu and was expected to get better each day. I would describe any little changes that I had noticed. Our conversations lessened the tension I felt, helped me to get through another day, and kept me working with Tariq to help him develop. They couldn’t, however, wipe away the tears or bandage the wounds; that has been hard for them as well as for me.

My grandparents, who were still alive when Tariq was younger, would reassure me, too. I have a cousin my own age who has mild mental retardation and whose speech came late, and my grandparents helped my aunt and uncle a lot when Gregory was a little boy. They reassured me, based on how my cousin had developed. Even though he still stutters, he learned to read and write a little bit. He has a driver’s license and holds a civil service job as a janitor. All in all, he has done very well for someone with his ability. My grandparents seemed confident that Tariq would be like Gregory. Because of their direct experience and their love for me, I trusted their wisdom and hoped desperately that they were right.

I remember how my grandfather would put Tariq on his lap and recite Italian nursery rhymes, the same ones that he had recited to me when I was a small boy. He would take Tariq’s hands within his strong palms, as he had taken mine, and rub them on his coarse beard until it tickled. Tariq would smile and then squirm to get away because he didn’t like to be held for long. My grandmother, too, would try to hold him, and she would sing the lullabies that she had sung to me, my siblings, and my cousins when we were little.

From Robert Naseef:

“Because autism affects communication and behavior, it strikes at the core of your relationship with your child. It is vital to get intensive and individualized services as soon as possible. Other parents of children with autism may be your best sources of information and support, so I always urge people to take advantage of support groups.

“I also stress the importance of accepting where your child is on the developmental ladder. Don’t push all the time, follow your child’s lead, and find mutually enjoyable activities. Your relationship with your child along with appropriate interventions and therapies will help your child grow. Spending some time each day just enjoying your child is also great chicken soup for the parent’s soul.”

No parent wants his or her child to be noticed as different from the norm. You want a healthy, happy child who fits in and makes you proud. But what do you do when your...
child is different? Speaking for myself, I had a hard time when my son was diagnosed with autism. I walked around looking out of the corner of my eye ready to pounce on people for their reactions. Actually, it was more my perceptions of their reactions, not to mention the perpetual knot in the pit of my stomach. Like many parents whose children are different, it felt to me that I was battling the whole world.

I wish we had Everybody’s Different: Understanding and Changing Our Reactions to Disabilities by Nancy Miller and Catherine Sammons much sooner. These authors give us an enlightening look at the many differences that exist among people in every walk of life and explain that these differences need not divide us socially. Life is definitely easier when you understand your own reactions, as well as those of others, while realizing that you are not alone—not by a long stretch. The authors help us understand our internal emotional reactions to disabilities and the differences that result from those conditions. They teach us strategies to build our lives, our families, and our communities in a way that is inclusive and respectful of our differences.

When Tariq starts “stimming” (i.e., repetitive behavior patterns that are a prominent characteristic of children and adults with an autistic spectrum disorder) or flapping, especially now that he is full grown, I automatically want him to stop and act “normal.” Internally, my reaction can be rather desperate because I don’t want him and me to be noticed as different. I also still find myself trying to avoid taking photographs when he is not behaving “normally.” There is a certain image that I prefer of him in my head, as much as I hate to admit that. But his behaviors are, after all, a part of him.

Even though I would consider myself a veteran parent, I have struggled to understand and own up to these reactions. It has taken me some time, but eventually I am learning to love my son not despite his differences—but rather because of them. The knot in my gut is long gone. Now, for the most part, I can look comfortably around me. I can notice what must have been there before—the friendly faces and kind eyes of compassionate folks.

I know I am not alone. Many parents have told me how isolated and abandoned they have felt in their hour of greatest need. There is a longing for someone to reach out and take a special interest in your child and therefore in you. When you finally do find that kind of support—a gentle hand to touch you and your child and a kind heart to ask how things are—that’s when you know for sure that you are not alone.
Importance of Early Screening

In the 1980s, many children with autism did not have their first contact with a professional until they were school age. By definition, the characteristics of autism spectrum disorder appear before the child is 3 years old. Diagnosis at an early age will allow the design of personalized interventions that help maximize the child’s abilities. Early diagnosis also facilitates better communication between parents and child, better defines a child’s development, and the appropriate help parents would need to seek.

Early interventionists know the importance of developmental screening to identify disabilities or delays early in a child’s life. Nancy Wiseman is working hard to make widespread developmental screening a reality. Since 1998, she has served as founder and president of First Signs, an advocacy group committed to promoting developmental screening for all children.

At 23 months, Nancy’s daughter was finally referred for a speech and hearing assessment, and it was discovered that her receptive and expressive language skills and social-emotional development lagged far behind the norms for her age. She was eventually diagnosed with pervasive developmental disorder-not otherwise specified (PDD-NOS).

Ms. Wiseman happily reports that her daughter responded very well to intervention and now attends school with typically developing children. Although she still has some language delays and auditory processing difficulties, her cognitive and receptive skills are actually above the norms for her age. But, Ms. Wiseman adds, “we lost about a year [of potential intervention].”

In April of 2001, First Signs launched a pilot project in New Jersey. Working in collaboration with the Robert Wood Johnson Medical School, First Signs sent a mailing to every pediatrician in the state, offering them the opportunity to earn continuing education credits by participating in the First Signs program. The program consists of viewing the First Signs video on screening and developmental delays; filling out an evaluation questionnaire; and ordering the First Signs kit of screening tool samples, which also includes directions for their use, a milestones wall chart, and an early intervention referral guide.

First Signs continues to receive requests from pediatricians, parents, and other clinicians for information on early detection. It has also launched a mini-pilot program in Tallahassee, Florida, and is starting a statewide program, similar to the first started in New Jersey, in Minnesota. After finalizing a program model, First Signs would like to take the program nationwide rolling it out on a state-by-state basis.

The Assessment

Once a screening indicates signs of autism, the next step is to determine what type of assessment to conduct. A proper assessment will help to design an intervention program relevant to the child’s unique learning style and will provide parents with answers without misleading them.

The book LINKing Assessment and Early Intervention is designed to help intervention programs select an appropriate tool for assessing possible developmental delays. Dr. Stephen J. Bagnato, one of the co-authors of this book, gave us a brief overview of eight standards that program staff can use to evaluate assessment tools:
1. **Authenticity**—A good assessment/intervention system should encourage staff to observe the child’s behavior in natural settings. “Table-top testing” is not developmentally appropriate. Examples include a child’s ability to recognize a favorite toy or a McDonald’s sign.

2. **Collaboration**—The assessment method should ensure that parents and professionals are partners in decision making.

3. **Equity**—An assessment tool should allow staff to make accommodations for children with special sensory, motor, cultural, or other needs. The tool should measure a child’s accomplishments rather than require the child to have specific sensory or motor skills.

4. **Sensitivity**—There should be a sufficient number of items for assessing behaviors of the lowest-functioning children as well as the most capable. Each child’s individual strengths and needs should be identified.

5. **Congruence**—The method should be designed for use with children with disabilities; in other words, children with disabilities should be included in the normed group.

6. **Convergence**—The assessment approach should allow you to collect information from multiple people in multiple settings, including the home, and should incorporate both observation and interview data so that a professional is able to see the full range of a child’s abilities.

7. **Acceptability**—The assessment approach should be acceptable to both parents and professionals. The tool should allow for a socially acceptable approach, such as play-based assessment, instead of an IQ test. The process should tap “real” skills, such as a child’s ability to communicate with others, as opposed to a child’s ability to identify pictures.

8. **Utility**—The method has to be useful for a specific purpose, such as intervention planning. For example, an IQ test would be useless for linking assessment and intervention because it has nothing to do with a child’s development.

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From Dr. Bagnato:

“An assessment’s purpose is to create individualized plans,” notes Dr. Bagnato. “Goals then become benchmarks for measuring the progress a child is making.” It is essential, says Dr. Bagnato, that assessment, intervention, and progress evaluation be linked together. “It’s a check-and-balance system.”
Barry M. Prizant, Ph.D., CCC-SLP, Amy M. Wetherby, Ph.D., CCC-SLP, Emily Rubin, M.S., CCC-SLP, Amy B. Coggeshall Laurent, OTR/L, and Patrick J. Rydell, Ed.D., CCC-SLP, are collaborators on the forthcoming The SCERTS Model: Enhancing Communication and Socioemotional Abilities of Children with Autism Spectrum Disorder. They have developed a new approach to teaching children with ASD that addresses the three core challenges in teaching children on the autism spectrum: social communication, emotional regulation, and transactional supports. To share their work and findings, Rubin and Laurent spoke with us:

1. What gaps do you see in other educational practices developed for children with ASD that the SCERTS Model is able to successfully address?

In the SCERTS model, it is recognized that most learning in childhood occurs in the social context of daily activities. Therefore, efforts to support a child’s development occur with caregivers and familiar partners in everyday routines in a variety of social situations in the home, school, and community settings. Because a number of current educational approaches focus on working with a child in isolation of these settings and social partners, a gap in the provision of support for that child’s partners (e.g., family training, peer training, and support to service providers) is often created.

In the SCERTS Model, developmental objectives for the child in Social Communication (SC) and Emotional Regulation (ER) are addressed not only by focusing on the child, but also by measuring the progress of that child’s partners in their ability to implement Transactional Supports (TS) (e.g., interpersonal supports, learning supports, and environmental modifications) to achieve more successful social exchanges.

In the forthcoming SCERTS Manual, a curriculum-based assessment will be accessible for families and educators to gauge both a child’s development in social communication and emotional regulation as well as the success of that child’s partners in implementing transactional supports.

2. In what settings is the SCERTS Model designed to be followed?

As alluded to in the response to the first question, the SCERTS Model is designed to be a comprehensive framework that supports a child’s development of Social Communicative (SC) and Emotional Regulatory (ER) abilities. Within the SCERTS Model, the facilitation of these core capacities through the implementation of Transactional Supports (TS) is viewed as essential for supporting a child’s active learning. Since the SCERTS Model is based upon typical child development and social learning theories, it is critical that developmental objectives for SC and ER be addressed across a child’s educational, home, and community settings in order to successfully promote the child’s generalization of learning within natural activities and functional routines.

The overarching goal of the SCERTS Model is to enable children to participate more successfully in developmentally appropriate activities with family members, adult partners, and peers in a variety of settings. Therefore, to ensure social success from the onset of intervention, transactional supports must be implemented not just across settings, but also across all partners within a child’s life. The model works best as a team approach, as the expertise of professionals from different disciplines in partnership with families is viewed within the SCERTS model as the most effective and efficient way to best support a child’s development across a range of settings and partners.
3. How can the SCERTS Model be implemented in an inclusive classroom without detracting from the needs of typical children while providing individualized attention to students with ASD?

Within the SCERTS model, a child’s individualized education plan and educational placement are based upon many factors. First and foremost, a child’s individual strengths and needs are considered with respect to his or her social-communicative abilities and emotional regulatory capacities. In addition, the child’s family’s priorities, as well as the demands and resources of the available educational programming options are considered.

Regardless of the child’s educational placement, individualized Transactional Supports (TS) aimed at facilitating the child’s attainment and maintenance of a well-regulated state, a state which supports a child’s ability to learn, attend and “fit in,” are essential components of the child’s educational plan. Within an inclusive classroom, these supports may consist of both Interpersonal Supports (e.g., communicative style adjustments) and Learning Supports (e.g., environmental and curricular modifications). The implementation of these types of individualized supports within the natural routines and activities of a classroom fosters the child’s attainment of developmental objectives in the areas of Mutual Regulation and Self-Regulation. The child’s resultant expanding repertoire of regulatory abilities enables the child to be a more active and engaged member in the inclusive classroom setting. As an active participant in the classroom activities and routines, the child is an integral part of the classroom community, enhancing rather than detracting from the learning environment.

Likewise, many of the Transactional Supports (TS) implemented by partners, such as environmental accommodations (e.g., reducing extraneous stimuli) and other Learning Supports (e.g., schedules and visual organizers) may be of benefit to many of the child’s peers within the inclusive classroom.

4. Do you feel it is most beneficial for children with more severe cases of ASD to be taught in an inclusive classroom or special education classes? Why?

In the SCERTS Model, an appropriate educational setting is not a “prescription” based upon the level of “severity” of the child with ASD. Determination of social learning contexts must be a flexible process that is dependent on a child’s learning strengths and needs, the family’s priorities, and the demands of the child’s natural routines and ongoing educational programming.

The SCERTS Model provides a framework for determining natural social activities that are the most conducive to addressing educational priorities such supporting the most functional abilities in social communication and emotional regulation, goals that directly address family priorities, and goals that are developmentally appropriate. Since participation in one learning context, either entirely inclusive or entirely self-contained, may not fully address these priorities, it is more common that a child will be participating in a range of social learning settings throughout his or her day.

Having said this, interactions with children without special needs and children who could provide good social and language models in more natural activities and routines are viewed as an essential part of supporting communication and emotional regulation for children participating in a SCERTS Model program. In these more “natural” learning contexts, there are numerous opportunities for developing functional communication and social problem solving skills. These opportunities may also help “typical” children to become more sensitive and supportive partners by developing a greater understanding of how to
adapt to and develop relationships with children with developmental differences, thus promoting growth in both the child with ASD as well as the “typical children.”

5. One of the three main areas the SCERTS Model addresses, and that children with ASD typically have the most difficulty with, is social communication. What do you see as the key components for forming a relationship with a child with ASD?

In the SCERTS model, the social communication dimension has been designed to address the overriding goals objectives that help a child to become an increasingly competent, confident and active participant in social activities. These developmental objectives fall under two primary domains, the capacity for joint attention and the capacity for symbol use, recognizing that in addition to symbolic language and play, a child needs to develop strategies for establishing shared attention with his or her partners to ensure successful and mutually satisfying social exchanges and the development of relationships. Since becoming a competent communicator, however, relies on the support of a responsive social partner, the SCERTS model recognizes that direct instruction focused on the child in socially isolated “teaching programs” does not necessarily ensure positive social communicative exchanges and development of relationships across a number of different partners.

Therefore, to ensure social success, supports must be fostered across all partners and settings within a child’s life, including family members at home, same-aged peers and teachers at school, and those within the larger community. In the SCERTS Model, partner objectives are written to address Interpersonal Support modifications (e.g., communicative style adjustments) and Learning Support modifications (e.g., the use of visual modalities and environmental modifications). Relationships develop when both the child and the partner bear some responsibility for change and mutually enjoyable and successful exchanges are achieved.

6. Do you view stimming as a problematic behavior that educators and parents should try to eliminate from a child’s behaviors or do you see it as a positive behavior that falls under self-regulation, one of the goals of the SCERTS Model?

Behaviors that may be viewed by some as “self-stimulatory,” are often regarded as Behavioral Strategies for Self-Regulation within the SCERTS Model. Behavioral Strategies develop initially during infancy and are simple motor actions or sensory motor strategies that a child engages in to regulate his or her arousal level, remain alert, and/or self soothe. The use of Behavioral Strategies for the purpose of regulation persists throughout the course of a child’s development and may either be biologically driven (e.g., sucking one’s thumb to self soothe) or modeled by responsive partners (e.g., stomping one’s feet when mad).

Often times, children with ASD have difficulty responding to conventional Behavioral Strategies modeled by partners (e.g., squeezing hands together, tapping feet, chewing gum, etc.); and, therefore, frequently employ unconventional patterns of behavior (e.g., jumping, walking on toes, flapping hands, and fleeing from the social setting) regardless of their age. This is particularly true for children with ASD when they are experiencing periods of extreme dysregulation. In the SCERTS Model, these idiosyncratic behaviors are viewed as either “starting points” for attempts at regulation or as signs of dysregulation, not necessarily as “problem behaviors.” As a result, these Behavioral Strategies as well as the environmental and social factors associated with their display are considered carefully when designing Transactional Supports (TS) and targeting developmental objectives in the domain of Emotional Regulation.
Paula Kluth, author of You’re Going to Love this Kid! Teaching Students with Autism in the Inclusive Classroom, has been working with students with autism for many years. The book is a great insight for teachers of students with autism in inclusive classrooms and is filled with activities to build a strong community among the students in any classroom environment. We were lucky enough to get some direct feedback from Paula and we anticipate the arrival of her new book.

**You have a terrific ability to recognize the strengths of children with autism spectrum disorder. How important do you think it is to get students in the classroom directly involved with students with ASD?**

It is so important! Any teacher who wants a classroom that makes all learners feel safe, comfortable, and capable should think seriously about how to create a space and a curriculum that gives learners opportunities to share with and support each other.

For most students, the best part about coming to school is seeing and spending time with friends. For students who struggle to make social connections, however, going to school can be a lonely and frustrating experience. Many students with autism who are being educated alongside their peers without identified disabilities are indicating that they need more than an inclusive classroom to feel successful; students with autism are increasingly asking teachers to facilitate the development of friendships and of access to social opportunities.

While no teacher can create friendships between students, every educator can create conditions in the classroom that will give students opportunities to strengthen social relationships, learn about/from each other, and get/give support—the hope being, of course, that these opportunities eventually lead to the development of friendships.

**How do you think inclusion affects the learning capabilities of students with ASD?**

An inclusive education gives students opportunities to access academic content, study social interactions, and learn new skills and information from a variety of sources (teachers, peers with a range of abilities and talents, classroom materials).

Students with autism often profit from watching and interacting with others their own age. Since many of these learners are master imitators, watching 20 other students learn to read or use science lab equipment can be more powerful than watching the teacher or following written directions. Students in these classrooms typically serve as natural supports for one another; they serve as models for speaking, listening, reading, writing, and using new materials. They also teach new skills to each other as they play, socialize, and work together.

**You have said in the past that sometimes the challenge of inclusion is more about the classroom than it is the kid. What would be the first steps for teachers in setting up their classroom for inclusion for the first time?**

Yes, in some instances, the teacher is doing everything right—effective teaching strategies are in place, instruction is motivating and responsive, and the curricular adaptations are all appropriate, but the environment is in some way problematic.

For instance, some students with autism struggle with certain sounds. Some of the sounds that disturb students might be obvious. Many learners with autism, for example, will scream when the fire alarm sounds. Some students with autism report, however, that they are equally bothered by some “everyday” sounds like buzzing electric pencil sharpeners and whirring air conditioners. To help students deal with classroom sounds a teacher might try earplugs or headphones for some activities or for use in some parts of the school building.
(e.g., gymnasium); reduce classroom noise when possible (e.g., have students push chairs in quietly or put carpet samples under chairs); or allow students to listen to soft music using headsets in noisy or chaotic environments.

Some individuals with autism also have incredible sensitivity to light. Children and adults with autism have reported problems in particular with florescent lights. The teacher may want to turn off the overhead lights for a few days to see if change seems to have an impact on the student.

Finally, appropriate seating may not be the first thing a teacher considers when planning for a student with autism, but for some students comfortable classroom furniture is pivotal to their success. One of my former students couldn't sit in a desk for more than a few minutes but he could sit in a beanbag chair for 40 minutes at a time. We soon purchased several beanbag chairs for the school (a few for the library, two for the music room, a handful for hallways, and one for the office) so that this student could be at ease throughout the school and so that all students could enjoy a change in seating now and then.

**Your book focuses on students with ASD, but would you suggest teachers use the many activities with other students in the classroom?**

Of course. Since teaching is a dynamic process, the strategies that may work for one student may or may not work for another. Having expressed that, it is also true that there are certain strategies that are often successful with some students with autism as well as many students without identified disabilities. Many of these strategies, in fact, are simply suggestions for good teaching that can support students with a range of needs, strengths, and abilities in inclusive classrooms.

Students with the most significant needs often cause teachers to get creative and to stretch their idea of what it means to reach all learners. While a teacher can meet the needs of all students without ever having a student with disabilities in her classroom, these students are often a catalyst for change and creativity. Specifically, including students with autism may help teachers think more carefully about the choices students have, the design of the lesson, the ways in which students participate in teaching and learning, and the comfort, engagement, and opportunities for ALL.

From Paula Kluth:

“This book provides concrete examples of how to plan lessons, engineer a safe and comfortable classroom, provide communication opportunities, and understand and support challenging behaviors. Drawing on classroom and school observations, as well as my own experience as an elementary and high school teacher, I explore pragmatic ways of making schools safe, challenging, and accessible for students with autism spectrum disorders.”

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Teachers who are good listeners often find that students are able to provide them an exceptional education about teaching, learning, and dozens of other topics. One way to listen is by tuning into students as they work and socialize. Teachers who participate in casual conversations at lunch time or “hang out” in hallways can learn about students in more holistic ways. What are students excited about? Worried about? Knowledgeable about? Interested in? Afraid of?

Teachers may find keys to teaching and learning and discover ways to better support students by listening carefully and respectfully during informal, but important moments. One teacher found ways to listen to her students by forming an “anyone-is-welcome” poetry club. She benefited as a teacher not only by participating in a social activity with students, but by eliciting their voices through the poems they wrote.

Educators must also be constantly scouting for students’ talents and seeking situations that highlight their abilities and support the needs of diverse learners. Teachers of students with autism must believe that students are competent and then they must “set the stage” for students to perform competently. Teachers looking for competence and complexity in learners should constantly consider questions such as Under what circumstances does this student thrive?; What gifts/skills/abilities does this student have?; How can I help this student find success?; and How and what can I learn from this student?
C. Organization

Possessions 1. Recognizes personal belongings (mine)

**Strategies**

- Make distinct the separation between the child’s possessions and others’ by
  - Marking the child’s space and materials
  - Clearly defining community space and materials
  - Identifying shared space and materials

Designate spaces for personal belongings (e.g., coat hook, chair, cubby, shoe rack, table area, work materials) using the child’s photograph and name.

Create personal tags with the child’s name and photograph to label items the child owns or is using.

Define possession by color-coding objects in the environment. For example, all of the items with red circles belong to the child. Items with purple circles belong to peer A. Items with blue circles belong to the teacher. The child does not have free access to items not marked by red circles. Mark common materials with multicolored dots.

Give the child a box. Prior to free play time, have the child fill it with his toys of choice. All items in the box belong to the child. Any items not in the box belong to others.

Set up a joint tabletop activity (e.g., playdough, art project, puzzle, simple building task) with shared materials. Give each child a small bin. Divide the materials between the child and his play partner by filling their bins. Define for the child that the items in his bin are his and the items in the play partner’s bin belong to the play partner. If the child wants an item that is not his, prompt the child to identify who has it and direct his request to that person.

Give each child a place mat colored or labeled (with the child’s photograph and name) to indicate his personal play space. Present play materials in one centrally placed box. The items placed on the child’s mat are his, the items on the play partner’s mat belong to the play partner. Materials in the box are shared.

Use woven place mats with two colors to represent a shared play space.

Engage the child in show-and-tell activities. Have the child bring in a personal item to show and describe to his peers. When other children bring in items, have the child identify which items belong to whom.
Activity 2
Social Skills Curriculum
Social skills should help children with autism understand what to DO, who to WATCH, how to LISTEN, and what to SAY.

Activity
B. Group Skills
Following Group Directions 1. Nonverbal directions

Selecting Activities
When preparing the child for a group activity, consider the following:

- Does the child attend to the adult in the group?
- Does the child follow the nonverbal directions in a one-to-one situation?
- Does the child understand the nonverbal directions used in the group?
- Does the child follow the lead of peers when confused?

Sample Activity: Group instruction

Design a social story about following group directions. The story should describe one or more types of nonverbal directions and explain what to do. For example, it is time to stop and look at the teacher when the lights flick on and off. Read the story to the class several times a day. Silently flick the lights whenever it is time to give group directions. Wait for all of the children to stop what they are doing and look at you before giving any verbal directions. If the children have not stopped, have another adult silently stop them and point them in your direction. Fade the silent, physical prompting as soon as possible.

Strategies
Organizational Supports
- Turn the lights on and off.
- Ring a timer.
- Ring a bell.
- Snap to a beat.
- Have everyone raise their hands and put a finger to their lips.
- Clap to a rhythm.

Social Supports
- Use similar nonverbal group directions across the day.
- Use exaggerated gestures.
- Target a peer coach for the child to watch in order to remember what to do.
- Assign a peer partner to assist the child.

Visual Supports
- Make a visual list of the group rules: watch, wait, and listen. Cue the children as needed.
- Provide individual cue cards that specify the group rules.
- Preview and review a social story about group rules.
Activity 3
Communication Skills Curriculum
The activities here enhance three areas of communication: basic communicative functions, socioemotional skills, and basic conversational skills.

Activity
B. Socioemotional Skills
Express Feelings 1. Requests a break when upset

Sample Means

Communicative gestures: Moving person's hand, reaching, pulling person's hand
Emergent: Vocalizations or speech, signs, photographs, pictographs, a voice output system, or written language to say Break or other appropriate word
Verbal: Speech, signs, photographs, pictographs, a voice output system, or written language to convey a multword message, such as I need a break, please

Learn a Target Message to Replace Behavior(s)

Context: All settings in which the child demonstrates a high level of discomfort or frustration
Materials: A tangible symbol (e.g., object, photograph, pictograph, written message) that means break to the child—children with autism often lack the ability to verbalize any communication that is a replacement for problem behavior, so a tangible symbol is essential even for verbal children
Procedure: Work with the child in a setting that is challenging. Wait for the child to demonstrate some level of agitation. Present the break symbol to the child and silently guide him to a clearly defined break area. A break should be calming for the individual child but not reinforcing. Maintain a neutral affect and do not engage the child in any discussion. Examples might include sitting on a beanbag chair or walking in the hall. The break must have a definitive end point and the child must come back to the activity. A request for a break must never equal a request for All done. Continue to anticipate the child's frustration and move from picking up the symbol yourself to gesturing toward the symbol and, finally, to the child initiating that process. The final goal is for the child to request a break spontaneously, without any cuing. Repeat the sequence many times in the same manner. Initially, all of the child's communication attempts should be acknowledged, even though he may overuse this new privilege. Gradually limit the number of breaks that are available. Once the child is requesting breaks from various adults, he can also be encouraged to tell a peer.

Strategies

- Teach meaningful social and communication replacement behaviors at the moment the child is engaged in inappropriate behavior.
- Offer the child maximum support to ensure success.
- Do not prompt in exactly the same way every time.
- Limit the use of verbal prompts in the form of questions.
- Simplify language complexity.
- Use AAC supports that are consistent across various settings.
- Keep the length of breaks consistent and clearly define the end point with a timer or clock.
- Observe what situations frustrate the child.
- Ignore any of the child's attempts to engage in conversation; remain neutral.
- Clarify expectations.
- Use nonverbal reinforcement that mirrors the child's emotional state.

Nick was a 12-year-old boy who had received a diagnosis of autism. He lived at home with his parents and a sister and attended a general sixth-grade class at his neighborhood school. Of most concern to his parents and school were his frequent self-injurious behaviors, which mostly involved his poking himself in the eyes. He also frequently would get upset at home and at school, throwing objects and sometimes even breaking windows.

We first had contact with Nick because of his very disruptive and disturbing behaviors. As we talked with his family, we found that he also had difficulty at bedtime and throughout the night, and we suspected that his sleep problems may be contributing to the daytime behavior problems observed at home and at school. One of the major problems surrounding Nick’s sleep was his unwillingness to go to bed at night. Often, he would remain awake for up to 4 or 5 hours after he was put to bed for the night. During this time, he frequently would scream, yell, throw things around his room, and bang on the walls with his fists. Unfortunately, this was almost an every-night event.

As happens very often in children with bedtime problems, Nick also sometimes had trouble remaining asleep at night. Several nights each month after he fell asleep, he would awaken, be unable to fall back asleep, and then become disruptive in the middle of the night. Two or three times each month, he would not sleep at all and would have tantrums on and off throughout the night.

Before we had contact with Nick and his family, a physician had prescribed several different medications to try to help him sleep. As is often the case, they tried giving him some Benadryl at bedtime to make him drowsy. Unfortunately, Benadryl seemed to have just the opposite effect on Nick, making him more agitated, and he stayed awake later than ever. For a short time they tried a drug called Inderal—which is used to control high blood pressure and migraine headaches and is also used sometimes to reduce anxiety—but this drug seemed to have no effect on his sleep. A third drug, Mellaril, was prescribed to be given right before bedtime, and this seemed to help Nick fall asleep more easily. The positive effects of using this drug were not without cost, however. During the first 4 months that Nick was taking Mellaril, he gained 40 pounds. One night he somehow got access to the bottle and drank a large amount of the drug and needed emergency treatment. Because of these concerns, Nick’s parents were anxious to try a different approach that did not involve the use of medication.

Nick’s parents had already established a fairly stable bedtime routine. When we looked at his sleep diaries, we found that, even with the Mellaril, it would take him an average of almost 2 hours each night for him to fall asleep. His parents had selected an 8:00 p.m. bedtime, in part because they wanted him to get more sleep and also because they relished time alone at night for themselves. Our suggestion for a later bedtime (10:00 p.m.), which was probably more appropriate for a child his age, was met with obvious resistance. The compromise that we all agreed to involved continuing to put Nick in his bedroom at 8:00 p.m., but instead of turning out the lights and trying to get him to sleep, he would be allowed to stay up and play in his room until 10:00 p.m.

From Mark Durand:

“A parent’s proper response to bedtime is essential. A 30-minute bedtime routine that is designed to prepare a child for sleep. Activities during this period can include reading to the child, washing, and getting undressed. Activities to avoid include any sort of aerobic activity (which increases body temperature and makes it harder to fall asleep) and any intake of caffeine. In fact, Dr. Durand recommended that children avoid exercise (including rough housing) and caffeine for a few hours before their bedtime.”

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Each evening, Nick’s parents would begin the bedtime routine, which would end with his being in his room at 8:00 p.m. He could keep his light on and play until 10:00 p.m. If he were very disruptive, then the parents could go into his room, but other than these times they were to leave him alone. An alarm clock was set to go off at 10:00 p.m. to signal both Nick and his parents that it was time to sleep. At this time they would enter his room and sit by his side for a few minutes of quiet activity (backrubs, quiet talking). After no more than 15 minutes, his parents were instructed to say goodnight, turn off the light, and leave the room.

The compromise at bedtime dramatically reduced the disruption at night. During most evenings, Nick did not fight bedtime and generally cooperated with going to sleep. There were still one or two nights each week when Nick was disruptive, however, so we designed a graduated extinction plan for these times. On nights when Nick refused to stay in his bed at 10:00 p.m. and go to sleep, his parents were instructed to wait 5 minutes before going into his room. We suggested that they stand by his closed bedroom door so that they could hear him and so that they could respond when he tried to leave his room. If he opened the door and tried to come out, then his parents would lead him back to his bed without saying anything other than, “Go back to bed.” On a few nights, they could hear him banging his head, so they calmly entered the room, placed him back into bed, and then left the room. Over the course of several weeks, the problems continued to decline, and Nick’s parents were delighted with the changes.

The solution of having Nick stay in his room from 8:00 p.m. until 10:00 p.m. obviously was a compromise that was not without some risk. By letting him spend so much time playing in bed at night, we were concerned that he might associate the bed with play rather than with sleep, and this might interfere with his sleep. Fortunately, this was not the case, and on most nights he fell asleep soon after the alarm went off. Ideally, his parents should have kept him up until 10:00 p.m., but the needs of the family—some “mental health” time together—were important to consider when we designed the plan. Its success was welcomed by the whole family.

Again, Nick’s case illustrates the need to tailor these programs for each family. When children present multiple problems surrounding sleep, as Nick did, it is important that you be patient and continue to monitor your child’s progress. Parents should complete the sleep diaries throughout the time of the program so that they can see whatever changes are occurring, even if progress is slow. Nick’s parents initially were skeptical about the program until we showed them his improvements each week. Seeing that Nick’s tantrums were becoming shorter and shorter gave them motivation to keep going. Remember to keep monitoring your child’s progress, and, if you need it, use this information to help you persist.
This case history involves an interdisciplinary evaluation of a 20-month-old boy known as Josh. Josh was referred to the interdisciplinary team by his primary physician due to parental concerns regarding Josh’s ability to communicate and difficulty gaining his attention. Josh was not receiving any intervention services at the time of assessment. His parents noted a great deal of variability in Josh’s ability to use gestures and words and to respond to others. At 18 months of age, Josh was sent for an audiology evaluation due to his difficulty responding to others, and results indicated hearing was within normal limits. Josh’s physician and family requested an interdisciplinary evaluation at this time to assess Josh’s development and to obtain a thorough medical evaluation to examine the possible biological basis for Josh’s presenting concerns.

In a telephone conversation, Josh’s mother reported that her son was the fourth child in his family. His oldest sibling was 12 years old and had autism and mental retardation. Josh’s parents were very knowledgeable regarding autism and involved in their older son’s intervention programs. Josh’s mother said she was watching her younger children’s early development carefully because of concerns about having another child with autism. She stated that Josh was able to engage in many age-appropriate activities; at times, he used words and gestures, responded to his name, and pointed. However, she observed Josh had some autistic-like characteristics, such as becoming overly focused on tasks, being nonresponsive to others’ attempts to get his attention, and tending toward solitary activities. She also believed Josh had sensitivities to sound and tactile stimulation, but he ate a variety of foods. Overall, his mother requested information for helping Josh develop language skills and for managing his tantrums.

Based on the preliminary information, the team decided that evaluation should include professionals from the fields of speech-language pathology, psychology, psychiatry, and occupational and physical therapy, with the psychologist as the service coordinator. The purpose of the assessment was to provide diagnostic impressions and to evaluate the findings so that recommendations could be made for treatment. In addition, because this child was so young, the team decided that speech-language pathology, occupational therapy, and psychology clinicians would observe Josh and his parents during play before a specific assessment plan was developed. No preassessment conference was needed, as no new information was provided prior to the assessment.

Josh was seen on 2 consecutive days of evaluation beginning at 8:30 A.M., with each team member assessing Josh for 2 hours. The team decided that Josh first needed to be observed in a familiar and minimally stressful context to ascertain his usual level of functioning, so they began with a play-based assessment. Ideally, this observation would allow time for Josh to become maximally comfortable. However, based on the constraints of scheduling, this observation lasted for approximately 20 minutes.

It was decided that the speech-language pathologist, the occupational therapist, and the psychologist would combine their play-based assessments because, in the early stages of development, social, adaptive, and communication skills are very difficult to separate into specific domains. For instance, joint attention can be viewed as a communication and a social skill. These team members were very interested in observing Josh’s response to specific prompts that encourage early social and communication skills.
The main component of the play-based assessment was observing free play with Josh’s mother. The playroom, which had a one-way mirror, was stocked with toys that were developmentally appropriate and would elicit various skills or autistic symptoms. For example, a puzzle and a shape sorter were chosen to observe visuospatial skills while blocks were used to assess construction skills in addition to fine motor skills. Cooking and serving toys were placed in the room near a doll to encourage pretend-play scripts, a telephone and a car were chosen for functional play, and a ball was used to examine possible reciprocal play skills. A couple of books, along with drawing materials, were used to assess Josh’s response to visual stimuli; his attention to letters, pictures, words, and their joint attention; and his fine motor skills. A small box was placed in the room that contained assorted sensory-based items to examine Josh’s affinity for or aversion to various sensory inputs. The box contained a Koosh ball, a textured hedgehog squeeze toy, a magic wand with floating stars inside of it, and a pom-pom. Also, large cushions were placed in the room to facilitate comfortable floor play. Often they are used by children and parents for hiding and tickling games or as a way to apply deep pressure to children who crawl under the cushions. Sometimes children have strong interests in items such as trains, cars, or particular characters. When this occurs, efforts are made to include these things in the assessment in order to observe the child’s level of interest and flexibility around them. However, Josh’s mother did not report that he had any fascinations other than letters.

Josh and his mother were asked to play together in the room with the mirror while the occupational therapist, speech-language pathologist, and psychologist observed from behind the mirror for 20 minutes. Josh’s skills were assessed in a number of areas of development, including typical and atypical developmental characteristics. After the first few minutes of the observation, it was clear that Josh was overwhelmed with the number of toys in the room. He randomly ran around the room, occasionally throwing the toys. After conferring with Josh’s mother, the decision was made to structure the playtime to a greater degree. His mother cleared the toys into a pile and systematically tried to engage Josh with them.

During the play-based evaluation, the clinicians observed Josh’s behavior in order to assess his social, communicative, motor, cognitive, play, organizational, and sensory development. These observations were placed within a general developmental context, so that a picture of Josh’s overall development could be obtained. (The reader is directed to Knox, 1997, and Linder, 1993, for descriptions of how to structure and interpret a general play-based assessment of young children.) Josh’s behavior was also monitored for atypical behaviors and those that are strongly associated with autism (see Table 14.1). To a lesser extent, Josh’s mother’s behavior was observed to assess the general quality of the mother–child relationship and the mother’s ability to read Josh’s cues and to provide support.

**Semistructured Play Interview**

After the unstructured 20-minute play observation, the speech-language pathologist and the psychologist developed a list of skills that had not been adequately observed during free play. Then attempts were made during the semistructured play interview to elicit them.

The speech-language pathologist and the psychologist worked together for 40 minutes, alternately entering the room with Josh and his mother in order to prompt for and examine specific social, communication, and play skills as well as his attention and repetitive motor behaviors. The majority of this interview involved items from the Autism Diagnostic Observation Schedule– Generic (ADOS-G; Lord, Rutter, & DiLavore, 1998). This is an empirically validated play-based diagnostic assessment of autism spectrum disorders. Ideally, a full ADOS-G would have been administered. However, due to time constraints, some portions of the ADOS-G had to be omitted. In addition to the ADOS-G elements, other
prompts were introduced such as examining Josh’s response to another person’s distress and the communication assessment prompts described by Wetherby and Prizant (1992, 1998).

**PROCESSES AND FINDINGS OF INDIVIDUAL DISCIPLINES**

After the initial play-based assessment and the semistructured interview, the psychologist and the speech-language pathologist evaluated Josh for an additional hour each. On the second day of testing, the occupational therapist and the developmental psychiatrist evaluated Josh for 2 hours each. The findings from these individual evaluations are explored next.

**Psychology**

The goals of the psychologist's evaluation involved assessing Josh's skills for diagnostic purposes, documenting eligibility for services, and recommending interventions. Standardized measures, both parent report and cognitive assessment, were to be used to ascertain the severity of Josh's delays compared to same-age peers and, in turn, to establish eligibility for services. The parent report of Josh's adaptive skills would give a good picture of his capabilities at home. Josh’s performance on nonverbal measures would provide information on Josh's cognitive abilities when language is minimized. (A full IQ evaluation would not be performed because of time limitations, likely compliance issues, and the fact that the speech-language pathologist would perform an assessment of Josh's expressive and receptive language.) Standardized testing would also allow observation of the child's ability to comply with instruction and his flexibility, and the examiner could begin to anticipate Josh’s initial response to more direct therapeutic interventions. Previously monitoring Josh in free play and semistructured play gave an opportunity for assessment of Josh’s typical social, communication, and play skills as well as his use of the environment. This observation also allowed the clinician to begin to anticipate how Josh would respond to more play-based approaches and to document Josh's development for treatment planning and in measuring future progress.

Across these assessments, the clinician would develop a picture of Josh’s overall cognitive abilities. Once this impression was formed, Josh’s nonverbal social and communication skills could be assessed in relation to his general nonverbal cognitive level for diagnostic purposes. Thus, if Josh displayed social skills at the same level as his nonverbal skills, a diagnosis of autism would not be considered.

**Assessment**

The first standardized assessment tool used was the Mullen Scales of Early Learning (Mullen, 1992). It is likely that Josh would be able to demonstrate higher abilities during incidental play at home when allowed to work on tasks on his own terms. However, given that Josh was able to comply with some of the testing demands, this evaluation yielded a reasonable estimate of Josh's abilities in a standardized testing situation. Two scales of the Mullen were administered to Josh in order to assess his visuospatial and visual-motor development. On the Visual Reception Scale, Josh's accomplishments were at the 18-month-old level, which placed his skills in the average range. On the Fine Motor Scale, Josh's performance was in the average range at the 18-month-old level.

The next standardized test was the Vineland Adaptive Behavior Scales (VABS; Sparrow, Balla, & Cicchetti, 1984). Based on parent report, Josh displayed delays in communication, daily living, and social skills. In the overall Communication category, he received a standard score of 69, with receptive and expressive age equivalents being 10 and 8 months respectively. Josh received standard scores of 69 in Daily Living Skills and Socialization. The Motor Scale was not performed due to time constraints and the occupational therapist’s use of parent report measures in her evaluation.
In addition to standardized assessment measures, nonstandardized techniques were used in this portion of Josh's evaluation. Play observations yielded a wealth of information. In terms of social development, Josh displayed a great deal of variation in his ability to attend and respond to his mother and the evaluator. He turned to the examiner when she entered the room but did not respond to the examiner calling his name at a later time. Josh did, however, respond to his mother the second time she said his name. In addition, before the majority of toys were removed from the room, Josh rarely attended or responded to his mother. After the toys were removed and his mother presented materials one by one, Josh took interest in his mother's actions when she was playing with toys that were enticing to him (e.g., books, puzzles). Once he obtained the appealing toy, the child tended not to look at his mother, although he did imitate her previous actions with these toys at times. As for imitation without objects, Josh clapped his hands to a game of Pat-a-cake but did not imitate any other actions, even when prompted. Josh was also able to engage in a social routine that involved tickling and swinging. He smiled and ran away when his mother introduced an “I’m gonna get you” game. After running away and hiding behind the chair, he turned and looked at her while smiling, waiting for her to catch him. Overall, however, Josh tended not to look at adults when smiling. He rarely alternated gaze between objects and adults, and this only occurred for the purpose of requesting. He did not point or show objects for the purpose of commenting, yet it appeared that Josh pointed at pictures in order to label them. When the evaluator feigned hurting her finger, Josh looked briefly at her and then returned to his activity. This suggested delays in his ability to notice and respond to emotional displays.

Josh revealed varying abilities to communicate. During free play, he did not use any intentional communication strategies. Most of the time, he ran around the room and produced repetitive vocalizations. When his mother and the evaluator presented toys in a systematic, routine, and highly structured manner, Josh was able to employ more sophisticated communication skills, especially when toys of optimal interest were used. For instance, Josh and the evaluator developed a routine in which the evaluator would hold the bubble wand up to her face and say “Ready, set . . .,” and Josh was able to state “Go” in order to request that the evaluator blow the bubbles. As the pattern became familiar, Josh sporadically combined eye contact with go. However, Josh was not able to engage in routines with objects if he was overly interested in them. At these times, it appeared that he had difficulty taking his eyes off or releasing possession of highly preferred objects, even when he wanted help with them. For instance, he became quickly upset when the wind-up car stopped but was unable to request help in order to get it to go again, even when given a high level of prompting and support. Josh used some repetitive vocalizations during this time as well. (See the speech-language report for a more thorough description of Josh’s communication skills.)

Josh’s toy play and use of the environment were also noted during the play observation. When too many objects were present in the room, Josh appeared overstimulated and his behavior became more disorganized and self-stimulatory in nature. Josh also tended to become focused on some toys to the extent that it was difficult to obtain his attention by calling his name. Nonetheless, if another highly interesting toy was presented, Josh was able to attend to this new object. In addition, when the toys were presented one at a time with no other objects available, Josh was able to play in a more organized fashion. Josh demonstrated some beginning functional play skills by rolling a car around the room and placing a telephone by his ear. Even when prompted by the examiner, Josh did not display any early pretend play, such as talking into the telephone or feeding dolls. Josh used some toys in a slightly repetitive fashion. He also tended to run around the room when activities were not
structured, which his mother reported often occurred when Josh was in new environments. Overall, however, the child did not perform any clear repetitive motor actions.

**Impressions and Commentary**

Based on observation and testing, Josh exhibited delays or atypical development in a number of areas. For example, he presented with impairments in social development. Specifically, these included lower than average responsivity and eye contact with others, difficulty showing or pointing to objects for the purpose of sharing his interest, and trouble understanding emotional displays in others. He also evidenced delays in his ability to use gestures to communicate and a lack of pretend play. At times, Josh was highly focused in play that had a slightly repetitive quality. Nevertheless, he could be easily distracted with another preferred toy, and he often repeated actions that had a functional purpose, such as using a shape sorter a number of times.

It should be noted that Josh had a number of considerable strengths. His nonverbal abilities were near age level, and he had an early interest in numbers and letters. Josh was able to use toys functionally on his own at home, particularly puzzles and shape sorters, and this boded well for Josh’s continued cognitive development. In addition, Josh was able to engage in interaction with adults when it was structured and routine. In these contexts, he demonstrated intentional and functional communication skills as well as the ability to share smiles with his mother. These qualities provided a good foundation upon which to build more complex social, cognitive, and communication skills.

Josh had a great deal of difficulty completing the cognitive assessment. He refused many items that his mother reported he could easily complete at home, and Josh displayed higher nonverbal skills during the free play when he could use toys on his own agenda. In this way, Josh’s assessment may have underestimated his abilities. However, the testing was important because it established that Josh’s nonverbal cognitive skills were at least near age level, ruling out a global developmental delay. In contrast, Josh demonstrated nonverbal communication and social skills that were scattered, with none higher than the 9-month-old level (e.g., use of gestures and alternating gaze for requesting, participation in ritualized games). This meant that Josh’s social and nonverbal communication skills were far below what would be expected for a child at his near-age level of development. If Josh had displayed nonverbal cognitive skills below a 9-month-old level, it is unlikely that a diagnosis of an autism spectrum disorder would be considered. This diagnosis would be discussed in the team meeting after the completion of all discipline evaluations. Because a full ADOS-G could not be completed, a diagnosis based on ADOS-G algorithm could not be computed. Nonetheless, many of the tasks from the ADOS-G gave the clinician important opportunities to observe Josh’s social and communicative behavior in prompted situations.

**Occupational Therapy**

The occupational therapy assessment was to include standardized testing, parent interview, structured play observations, and clinical observations. Standard performance scores would allow for easy identification of areas of concern. Gross and fine motor skills would be measured through standardized testing and clinical observations. Due to Josh’s high need for control, it was likely that standardized test scores would not reflect abilities that Josh displayed day to day. Therefore, the goal of standardized testing was to obtain scores that could be used to establish eligibility for services, to reflect Josh’s abilities relative to typically developing peers, and to provide a baseline for comparing future performance.
A structured parent interview would be used to gather information regarding self-care abilities and the functional use of fine motor skills. Interviewing could be completed while the child was playing or over the telephone if time did not permit an in-person interview. Play skills would be measured through two observation periods. First, team observations of Josh and his mother interacting with selected toys would reveal functional toy use, initiation, toy preferences, quality of movement, access to and manipulation of toys, and his mother’s ability to gauge and respond to Josh’s behavioral cues. Second, observation during occupational therapy assessment would include gross motor, vestibular, and sensory play experiences and focus on organizational skills, the influence of sensory experiences on organization and play, and experiences Josh sought or avoided. In addition, foundation skills such as muscle tone, movement patterns, range of motion, strength, endurance, protective responses, and soft neurological signs would be assessed through clinical observations.

Information regarding sensory processing abilities would be gathered through parent completion of a sensory history questionnaire and through structured observations. The sensory history questionnaire would be mailed to the family prior to the clinic appointment so that the therapist could review responses, identify any potential areas of difficulty or areas requiring further investigation, and develop a plan to evaluate the areas of poor processing in the clinic environment. Clinical observations would also involve watching for variations in response, indications of hyposensitivity or hypersensitivity to sensory stimuli, the impact of sensory experiences on arousal level, and Josh's ability to self-regulate his responses or the need for external support and assistance in modulation.

**Assessment**

Josh presented with hypotonia (low muscle tone) in his trunk and mild joint laxity in his hips and knees. He used a W-sit position for floor sitting. When placed in long sitting, his back was rounded and shoulders were slumped. Josh was fearful and clung to the therapist when placed on a therapy ball. He enjoyed slow, rhythmic bouncing but did not release his grasp of the therapist. He resisted lying on the ball. Ambulation was characterized by wide base of support and intermittent decreased dorsiflexion.

The Peabody Developmental Motor Scales (PDMS; Folio & Fewell, 1983) were utilized to assess gross and fine motor skill development. At times it was difficult to ascertain whether Josh was not capable of an activity or if he was merely resisting the requests of the therapist. Some modifications were made to standardized testing procedure in order to gain the highest level of compliance. For this reason, the scores reflected a fair estimate of Josh’s abilities in a standardized testing environment but may have been different than skills he displayed at home. On the Gross Motor Scales, Josh received a total score of 157, corresponding to the 2nd percentile, 2.05 standard deviations (SDs) below the mean, and an age equivalency of 16 months. On the Fine Motor Scales, Josh received an overall score of 129, corresponding to the 9th percentile, 1.38 SD below the mean, and an age equivalence of 17–18 months.

The Self-Care Domain of the Pediatric Evaluation of Disability Inventory (PEDI; Haley, Coster, Ludlow, Haltiwanger, & Andrellos, 1992) was used to guide a structured parent interview regarding Josh’s self-help abilities. Josh received a total score of 11, corresponding to a standard score of 22, a scaled score of 33, and an SD of more than 2.0, indicating delays in self-care abilities.

Josh’s parents completed the Sensory Profile (Dunn & Westman, 1995), on which they rated the frequency of several sensory-based behaviors. The findings were suggested sensory sensitivity. He demonstrated mixed responses to auditory and visual input, being sensitive to intense levels of input but appearing to not notice low levels of stimulation, particularly
when of a social nature (e.g., responding when his name was called). Josh turned to the
sound of the overhead paging system and often covered his ears in response to loud noise.
Josh also showed signs of tactile defensiveness, as his sensitivity to some textures (e.g., the
feathers in a duster) caused him to avoid a variety of play and learning opportunities. Yet this
sensitivity was not all-encompassing. Josh enjoyed pulling the strands of a Koosh ball and
playing with a prickly squeeze toy; he responded to the sound of the squeeze toy with laugh-
ter. Josh’s play skills were observed as part of team observations and in selected opportuni-
ties during occupational therapy assessment. Josh’s performance was partially interpreted
using the Revised Knox Preschool Play Scale (Knox, 1997). Results for the various domains
are summarized next.

In terms of motor skills, the child’s movements were functional but lacked fluidity, with
no rotation evident. Josh was able to obtain and manipulate toys, though the quality of his
movements was hindered by hypotonia. Josh enjoyed playing in the tunnel and sitting on
the trampoline while his mother jumped on it. He avoided all off-ground movement, refusing
the balance board and swing. When he was excited, Josh tensed his body and hands,
maintaining this posture for several seconds while intensely focusing on the intriguing activ-
ity. Josh was also noted to run in circles, producing vocalizations.

Josh’s attention and organization were limited. He was able to engage in play for more
than 5 minutes; however, in extended play, the quality became persistent. When fixated on
a toy, multiple cues and occasional physical assistance were required to help Josh shift his
attention. At times, Josh allowed his mother to join him in play, and he visually attended to
her toy use when she utilized his preferred toys.

Josh’s limited gross motor skills did not significantly affect his use of space in the assess-
ment room. He stood without support, walked with a wide stance, and threw a ball. His space
management skills were similar to those demonstrated by children 12–18 months of age.
Material management skills were also similar to those of children 12–18 months old. Josh
threw, inserted, carried, opened, and shut toys. He did not string beads, separate pop beads, or
use tools.

Josh’s toy play showed a preference for dump and fill activities, and he often repeated
this sequence. He poked his fingers into the holes of a pegboard but required assistance to
place pegs in the board. Josh stacked blocks vertically, pushed the buttons of a toy telephone,
and completed a shape sorter. He did not use markers.

Impressions and Commentary
Josh demonstrated some hypotonia, and gravitational insecurity was also evident. His hypo-
tonicity and weak abdominal musculature contributed to delays in gross motor skills and the
use of compensatory movement patterns. Fine motor skills were well within typical limits.
Self-care skills were delayed and influenced by sensory defensiveness.

Josh demonstrated fluctuating responses to auditory and visual input, suggesting diffi-
culty integrating information through these sensory systems. His hypersensitive responses to
movement and vestibular stimuli were consistent with gravitational insecurity, causing him
to avoid many activities involving movement. Josh’s negative reactions to unexpected touch
and avoidance of a variety of textures were indicative of tactile defensiveness.

Play skills were fair, as Josh demonstrated a predominance of skills typical of children 6
months younger than his chronological age. Hypotonia and gravitational insecurity impeded
engagement in functional motor play. Material management was a relative strength because
Josh demonstrated functional skills for manipulation of objects; however, he had a tendency
toward persistent use of objects. Inconsistency in imitation, interaction, and symbolic use of toys affected Josh’s ability to engage in functional play.

Minor procedural modifications were required for Josh to tolerate standardized testing. It appeared that his resistance to some items reflected his difficulty in tolerating imposed structure and transitions. Utilizing small toys and increasing the use of routine were helpful in increasing his cooperation, and these modifications to testing procedure were documented in the testing report. The play observations provided a snapshot of Josh’s abilities to gain access to play materials and apparatus and to manipulate tools and toys; ideation and flexibility in use of toys; participation in collaborative play; and skills in imaginative play, sensory exploration, and gross and fine motor activities.

Josh’s case depicts the importance of assessing sensory processing and the impact of poor sensory processing on arousal level. Completion and return of the sensory history questionnaire prior to the clinic appointment allowed the therapist to identify potential areas of lower processing. The therapist was then able to create opportunities during the testing situation to elicit some of the reported behaviors by including relevant toys and setting up the testing environment to contain significant sensory experiences. Thus, the quality and intensity of responses, as well as effective and ineffective methods of recovery, could be compared.

These play situations were important for observing the effect of sensory stimulation on Josh’s arousal level. The physiological state of Josh’s nervous system influenced how organized he could be in his interactions with the environment and the people and objects in it, as changes in sensory stimulation can cause fluctuations in arousal level. Therefore, it was crucial to assess each sensory system and determine which types of sensory stimuli caused such extreme fluctuations that he was unable to modulate his responses to the stimuli, resulting in behavior that was inappropriate for the situation. It was evident that Josh was easily overstimulated by the visual input of the toys scattered throughout the room. When the toys were gathered into one location, he was able to calm himself, leading to increased organization and the ability to engage in purposeful toy play.

On a similar note, it was necessary to assess the parent’s ability to interpret her child’s state of arousal and her effectiveness in helping him to modulate his responses to sensory stimuli. Most of the time, Josh’s mother was able to modify her style of interaction according to her son’s level of arousal. She decreased the level of auditory input by using short sentences, incorporated familiar games when Josh did not initiate functional play, and attempted to redirect his attention when he demonstrated signs of overexcitement.

Speech-Language Pathology
The aims of the speech-language assessment were to assist in determining a diagnosis, to document a need for services, and to develop appropriate recommendations for intervention to foster social-communication skills. Parent report would be
important for determining a diagnosis and skills and behaviors seen at home. Although Josh was very young, report of his development to this point (and any changes in development) would be needed for diagnosis. A description of current skills in a more familiar environment would be compared to skills and behaviors observed during testing in a less familiar environment. Standardized testing of language comprehension and expressive language skills would offer a standardized score or age equivalency and an opportunity to observe Josh’s interaction with materials and tasks that could not be modified. Semistructured tasks and observation of play with his mother would be used to assess and observe auditory attention and comprehension of language with varying levels of contextual, gestural, and visual supports. These settings would also be employed to ascertain expressive communication across the continuum from physical behaviors and vocalizations to gestures and words. Communicative intent would be evaluated along with other nonverbal social skills. Speech development, generally assessed through nonstandardized approaches at this age, could be observed in spontaneous as well as imitative interactions.

Assessment

Although many of Josh’s social and communication skills were exhibited during the observed sessions, additional information was needed to complete the assessment of communication skills. A standardized test of language skills, the Preschool Language Scale–3 (PLS-3; Zimmerman, Steiner, & Pond, 1992), was attempted but could not be completed due to noncompliance.

The play assessment yielded greater results. Based on earlier observations, toys and materials were brought into the room in closed, opaque containers and presented one at a time in small sets. Josh’s mother remained in the room and was asked to participate in some of the activities. A semistructured play-based assessment of language comprehension, expressive language, nonverbal social-communication, and speech-phonology skills was completed. Sensorimotor toys, thematically related toys, and toys that could be used to support social interactions were presented to elicit behaviors. The observations and behaviors obtained during the play-based assessments formed the basis for evaluations in the domains described next.

Language Comprehension

Josh exhibited decreased attention to auditory and verbal information. His focus increased when toys were presented individually into his visual field and when he was strongly motivated by the toy or activity. It was difficult to get his attention by calling his name when he was involved in a preferred activity. Josh showed no response to a loud sound outside in the hallway, and he generally required physical prompting to follow a direction. He also used context and routine to support comprehension of directions, sometimes making it unclear whether he understood key words in the verbal prompt or was using a strategy based on familiar routine. (For example, at home, he showed a consistent response to the words “bottle” and “up.”) Josh paused when the evaluator warned him, with a loud “no,” to stay away from an electrical outlet but continued to touch it.

Expressive Language

Josh appeared to be presymbolic, though he had been using some protowords (e.g., “da,” “ba”) and one-word approximations, such as “baba,” for bottle. Josh primarily vocalized a series of consonants and vowels that appeared to be self-stimulatory in nature. He produced one true word, “go,” in a verbal routine associated with a motor-based activity (e.g., “ready, set, go”) during the session. His mother reported he used this word consistently at home, and
“uh-oh” was heard sporadically at home. In addition, Josh’s mother stated that he would use a word a few times and then never use it again.

**Social-Communication**

Josh primarily demonstrated preintentional communication. He vocalized, fussied, or screamed to protest when toys were put away or when limits were set. He pointed proximally to pictures in books and to some objects but did not call the adult’s attention to the item in any way. These behaviors suggested that Josh may have been beginning to use a point to label. During the sessions, Josh was prompted to give the bubble wand to the evaluator to request more bubbles, but he resisted giving other objects when prompted. He seemed to lack the understanding that the object would be returned. Josh displayed reduced eye contact except when a motivating toy was held close to the evaluator’s eyes, and he glanced at her eyes when an action was delayed. He also rarely appeared to look at adults as he played with other toys. Josh protested and showed very early requests for the continuation of an appealing activity with significant structuring of the activity. He also used proximity to an object or container to signal interest in something. He attempted to get most things himself at home and was just beginning to take an adult’s hand and pull the adult toward a desired object.

**Speech Phonology**

Josh produced strings of consonants and vowels in a babbling-like sequence that appeared to be self-stimulatory. Consonants included b, d, and g. He showed some evidence of syllable formation in his protowords (“ba,” “da,” and sometimes “baba”) and one true word (“go”).

**Impressions and Commentary**

At age 20 months, Josh presented with significant delays in the development of language comprehension, expressive language, nonverbal social-communication, speech phonology, and play skills. He also exhibited atypical development in regard to his 1) decreased attention to auditory and verbal information, 2) difficulty shifting attention from a preferred activity to the source of a voice, 3) reduced eye contact and number of social-communicative interactions, 4) very limited use of conventional gestures to communicate intent, and 5) extended period of babbling with the production of self-stimulatory sound sequences.

Standardized assessment procedures could not be used to assess Josh’s skills for various reasons. First, testing was limited by materials that did not motivate Josh or hold his attention. Second, most receptive language items rely on the child’s motivation to follow directions. Given Josh’s high need for control, he found this test very frustrating. In addition, no modifications to the testing would engage Josh. The observations of play with his mother, along with the semistructured play-based assessment, offered the opportunity to use motivating toys and activities. It also allowed the examiner to observe Josh’s response to varying levels of support and prompting. The use of these approaches resulted in information that could be used to consider a diagnosis of an autism spectrum disorder. As this diagnosis was being considered, it was important to look at Josh’s atypical development as well as his delays.

**Psychiatry**

Much of the psychiatrist’s assessment was based on Josh’s past. The next section examines in detail information from different categories in the child’s history.

**Developmental History**

Josh was an “easy” baby who fell into a routine of eating and sleeping by his second month. Although Josh grew typically throughout his early life, since 15 months of age he had refused to eat meat. His mother had supplemented the child’s diet with multivitamins but Josh incon-
sistently took these as well. The parents noted some tactile and auditory sensitivities early on. Josh’s early motor development was slightly delayed but within typical limits. During infancy, Josh responded to face-to-face interaction at times, such as smiling when his parents spoke to him, and he engaged in physical baby games (especially if tickling was involved) by 10 months of age. However, it was more difficult to get Josh involved in nonphysical games such as Peekaboo.

Josh began using simple instrumental gestures, such as reaching his hands up to get out of his crib, by 12 months of age. At approximately 15 months of age, Josh began pointing to request foods that he wanted, and he started using a word to request actions. By approximately 18 months of age, Josh began pointing at objects that he obviously was not requesting, such as items in books. Yet he did not reference his parents when pointing; instead, his parents reported that this pointing appeared to have a more routinized quality. Josh would engage in rough-and-tumble play with his siblings, such as tickle games and chasing games. He did not display interest in peers, although he would engage in parallel play with a group of peers if the activity was motivating to him.

During periods of Josh’s intense interest, his mother noticed that he seemed even less responsive to his name. Although no one in the family ever observed any behavior that could clearly be called a seizure, these episodes of unresponsiveness, lasting 20–30 seconds, were a source of worry. There had been no loss of skills, vomiting, birthmarks, loss of vision, or visual impairment. Josh’s parents also reported their son’s sleep had been increasingly discontinuous at night. On some nights he slept no more than 5–6 hours, even though, with naps, he slept about 8–10 hours per 24-hour period.

**Medical History**

Josh was born as the fourth child to a 36-year-old, healthy female after a 44-week pregnancy, which ended in a vertex vaginal delivery after an 18-hour uncomplicated labor. His birth weight was 7 pounds 11 ounces, and he was 20½ inches in length. His Apgar scores were approximately 8 or 9. Josh went home and was believed to be well at 2 days of age.

Growth and developmental milestones were smiling at 1 month of age, rolling at 4½ months old, sitting at 6½ months of age, crawling at 8 months old, and walking at 14 months of age. Josh produced a few words, which he used sporadically, by the time he was 15 months old. Toilet training had not occurred at the time of this assessment. Tantrums were present.

There were no indications of heart disease, hypertension, diabetes, convulsions, strokes, meningitis, encephalitis, brain injuries, or abuse. Two bouts of otitis media resolved easily with antibiotics. No past surgeries, hospitalizations, allergies, or current medications were noted, and immunizations were up to date.

**Social History**

Josh lived with his parents and three siblings. His 12-year-old brother had autism and Josh’s two sisters had average or above-average intellectual abilities, although one had a mild learning disability. Josh’s family had private insurance, and their eldest son received Division of Developmental Disabilities (DDD) respite services. The mother was trying to obtain Supplemental Security Insurance for her older son. The parents described their relationship with each other as good, but it was strained at times by stresses involving their eldest son’s problems. Consequently, Josh’s parents had sought counseling once in the past and found it very helpful. They stated that they moved to the area 4 years ago and that they had good social supports through their church. Nevertheless, Josh’s parents reported being very concerned about his development. The mother, who was not employed outside of the home, was experiencing a high level of stress; the father, although concerned, seemed less stressed.
Besides an exploration of Josh’s history, certain on-site evaluations provided pertinent information. One of these was a mental status examination. Josh presented as a healthy child with no obvious dysmorphic features. He made repetitive utterances but used no discernible words. During the interview with his parents, he occasionally looked up but showed no distress when his parents left him alone with the interviewer. Instead, he played intensely with a brightly colored toy he brought from home. In fact, the child also was extremely difficult to engage and would not accept other toys offered him during the evaluation. At the end of the interview, when his plastic toy was put aside, he cried inconsolably. Aside from that episode, which was quickly resolved when the toy was given back, Josh appeared generally cheerful. There seemed to be no evidence of hallucinatory experience or staring spells, but he did exhibit other autistic-like characteristics. For instance, he withdrew from touches on the arm to get his attention. Furthermore, at one point in the interview, he climbed a chair to a paper towel dispenser in the exam room and repeatedly pulled out paper towels. When redirected, he ran around the room looking for his plastic toy.

A physical examination showed that Josh was at the 25th percentile for height and weight but at the 95th percentile for head circumference; these measurements were consistent with previous measures obtained by his family physician. His vital signs were normal. There were no birthmarks on his skin. Some of his physical features were marginally atypical (e.g., clinodactyly; small, slightly rotated ears; a high-arched palate). There were no cardiac murmurs or other physical findings. Josh’s neurological examination revealed no focal difficulties, but he had trouble running and jumping and tended to fall, although to no consistent side.

Impressions and Commentary
This 20-month-old boy presented with delays in language acquisition, gross motor development, and social behaviors. In addition, he demonstrated some atypical behaviors, including self-directed behaviors, developmentally inappropriate activity, and the inability to regulate attention. His mental status examination corroborated these historical findings as well as the fact that his play was nonsymbolic and would be repetitive at times. His sensorium was clear, and he did not evidence a psychotic process. Although he did not appear to have staring spells, this feature as the result of seizures had to be ruled out. At this time, the DSM-IV Axis 3 had to be tested to eliminate a diagnosis of fragile X syndrome, some other karyotypic irregularity, epilepsy, or an inborn error of metabolism. Also, a secondary nutritional deficiency from his restricted diet needed to be considered. On Axis 4, he probably was experiencing mild stress from his parents’ hypervigilance and marital difficulties. On Axis 5, he was globally functioning with mild to moderate impairments. This level of functioning was consistent over the previous year. Axis 1 and 2 DSM-IV diagnoses would be postponed until the multidisciplinary team meeting, once all the discipline evaluations had been completed.

Research shows a strong genetic component in autism is indicated by an increased concordance rate in monozygotic versus dizygotic twins (Bailey et al., 1995), and the risk to siblings of idiopathic cases is 75 times greater than that of the general population (Bolton et al., 1994). According to Folstein and Rutter (1977), about 2% of siblings are affected, and speech-language delay is common among siblings of children with autism. A later review by Smalley (1997) found that the sibling recurrence risk is 3%–5%. The implications of these statistics would be discussed with Josh’s parents, as it was likely that a strong genetic component for autism existed in this family. This point was upheld by the fact that Josh appeared to have and his brother did have an autism spectrum disorder; also, one sister had a learning disability.
Diagnostic tests were ordered to test for conditions based on symptom presentation and those known to co-occur with autism. A complete blood count (CBC)/differential test was ordered because of Josh’s restricted intake of meat proteins. If this lab test indicated iron deficiency anemia, Josh would be prescribed iron supplements in addition to a multivitamin. Karyotype was ordered to screen for any obvious chromosomal abnormalities, while DNA was examined for X-linked mental retardation syndromes such as fragile X. This test was necessary because the signs of fragile X syndrome in early childhood are nonspecific; developmental delay is an almost universal manifestation. Thus, any child (male or female) with a speech-language or motor delay of unknown etiology should be evaluated for fragile X syndrome, particularly in the presence of a positive family history and a consistent physical and behavioral phenotype and the absence of especially atypical structural features (Curry, Stevenson, & Aughton, 1997). When fragile X DNA-based testing is used regularly in this large and loosely defined group of children, the yield of positive test results is relatively low; approximately 3%–6% of unselected males have mental retardation (Curry et al., 1997). Urinalysis and an inborn errors of metabolism (IEM) screen were ordered to rule out phenylketonuria (PKU) and other metabolic disorders. The risk of these disorders is less than 1%; however, if present, they can significantly affect a child’s presentation.

**INTEGRATION OF DISCIPLINARY INFORMATION AND RECOMMENDATIONS**

The team met to discuss each member’s findings and to draw conclusions regarding diagnostic impressions and recommendations. Based on these evaluations, the following information was obtained in each of the areas:

1. A physical exam revealed no significant findings. Nonetheless, tests were ordered to rule out the possibility of any other medical conditions that could contribute to Josh’s overall delayed and atypical development. These results would be discussed with Josh’s parents when tests were completed.
2. Josh showed atypical development in the use of nonverbal social behaviors such as making eye contact, orienting and attending to social information, producing reciprocal interaction with adults, using joint attention techniques (e.g., pointing, showing) to share information, and understanding emotional displays.
3. In terms of communication, the child’s development was atypical regarding utilizing gestures, employing expressive and receptive language, engaging in pretend play skills, and vocalizing.
4. Delays in gross motor development and the ability to regulate sensory input in a number of domains impaired Josh’s capacity for organization and focusing attention.
5. The child’s attention span was limited; in fact, he possessed higher than average levels of distractibility and inability to focus when presented with competing stimuli. Josh also had the tendency to overfocus on some tasks.

**Diagnostic Process**

Given the preceding characteristics, a diagnosis of an autism spectrum disorder was likely indicated. Yet before confirming this diagnosis, the team discussed possible differential diagnoses. The psychologist’s testing showed that Josh’s nonverbal skills were within the average range. Ruling out a global developmental delay, the question arose whether Josh had an isolated language impairment. The assessment information, however, did not suggest this to be true. Josh displayed an inconsistent response to auditory and verbal information; he also had an extended period of babbling, and this babbling appeared to be self-stimulatory in nature.
In addition, the child’s development of communicative gestures was delayed, and the com-
 municative intentions expected by 20 months of age (especially “give to request” and point-
ing or showing to comment) were absent. The absence of complex and sequenced gestures implied that this was not a speech disorder, as most children with average nonverbal abilities and an absence of oral communication generally find a way to communicate nonverbally.

If Josh’s development remained consistent, his current profile suggested a diagnosis of a high-functioning autism spectrum disorder. Over time, a clearer picture would emerge regarding his specific diagnostic status. Because of his delays in language at the time of assessment, a late diagnosis of Asperger syndrome would be unlikely. However, at this point it was difficult to diagnose PDD not otherwise specified (NOS) versus autism due to issues of diagnostic instability at Josh’s young age. A future differential diagnosis would need to pay particular attention to the third category of behaviors that included Josh’s development of repetitive behaviors and restricted interests.

Recommendations
After the team deliberated on their individual findings, they jointly designed a set of inter-
vention recommendations. These items are described in the following section.

Educational Programming
Research suggests that early intervention can be highly effective for young children with autism when it is systematic and intense. The team strongly recommended that Josh receive early intervention services, with the majority of these services delivered in a one-to-one, struc-
tured, therapeutic context for the next year. His intervention program might also incorporate a few different approaches, including classroom time, speech-language therapy, occupational therapy, and individual therapy in a structured or play setting. Specific recommendations in each of these areas are One-to-One Intervention, Birth-to-3 Program, and Speech-Language Pathology and Occupational Therapy Consultation.

Speech-Language Pathology
The team recommended that Josh receive speech-language pathology services, in a clinical or birth-to-3 center, with emphasis on Language Comprehension, Expressive Language, Social-
Communication, Speech Phonology, and Play.

Occupational Therapy
Josh would benefit from clinical occupational therapy services to focus on gravitational inse-
curity, immature movement patterns, and self-care impairments. In addition, specific foci would strengthen this occupational therapy treatment, such as Sensory Processing, Physical Therapy Consultation, and Birth-to-3 Program.

Medical
Josh needed to be tested with a CBC, a karyotype and a DNA test for fragile X syndrome, gen-
eral urinalysis, and urinalysis for inborn errors of metabolism. A baseline of staring spells also needed to begin, and a consultation from a pediatric neurologist to rule out seizures was ordered. An electroencephalogram (EEG) and advanced brain imaging studies would be reconsidered after this consultation. The team would request follow-up appointments with Josh’s parents once the tests were completed to discuss the results and any further treatment.

Community Resources
Josh’s parents were already aware of the Autism Society, and it was likely that his mother would continue to find that group a helpful resource. In addition, Josh’s parents could also
contact the local Families for Early Autism Treatment (FEAT) group to obtain support in assembling and maintaining Josh’s home program. The clinic provided Josh’s parents with brochures from both organizations.

**Problem Behaviors**

It was likely that a number of Josh’s tantrums stemmed from either his frustration in not being able to communicate his needs or from his need for more structure. As Josh’s communication program became more systematized and incorporated into everyday situations, his tantrums would probably decrease. Furthermore, Josh appeared to become easily overstimulated by his environment, but he responded well and quickly to increased structure. The team members shared specific strategies with Josh’s parents for augmenting structure, and they included a handout on how to use visual supports (e.g., picture schedules, picture cue cards).

**Reevaluation**

The interdisciplinary team wanted to reassess Josh in approximately 1½ years to ascertain his levels of development and progress. In addition, his response to intervention would be assessed and subsequent treatment recommendations would be offered. A more specific diagnosis would probably be made at that time.

**Discussion with Parents**

A specific diagnosis (autism versus PDD or Asperger syndrome) at 20 months could not be established. Nevertheless, although a definite diagnosis was not provided for the parents, they were told that Josh clearly evidenced autistic characteristics. These features were described to the parents, including specific information regarding the severity of their son’s impairments. The parents were informed that although these characteristics would likely remain with Josh throughout his life, the team expected him to respond well to intervention. The clinicians also outlined Josh’s strengths, such as his ability and interest in interacting with others in structured predictable environments, his relatively strong nonverbal cognitive skills, his ability to engage in functional activities without parental support at home, and his emerging use of communication through words and gestures. In addition, the family’s strengths were noted: Josh’s parents were quite knowledgeable regarding autism spectrum disorders, and they were sensitive to Josh’s need for structure and predictability and his tendency for sensory overstimulation. All of these factors boded well for his future.

In the feedback session, the recommendations, as well as various intervention approaches, were discussed with Josh’s parents. Because his parents were already familiar with some interventions for children with autism, they were able to discuss the various approaches and the types of empirical and clinical support associated with each method. However, for many families that are first receiving a diagnosis of an autism spectrum disorder, specific information regarding intervention is often too overwhelming. In such cases, families need to be prepared for the wide array of interventions that are available for children with autism and the strong and often contradictory opinions regarding which types of interventions are most appropriate. One way to assist parents is to give them an overview of the major types of interventions that are available. Breaking them down into categories of those that have empirical support and those that do not can often be useful. (Freeman, 1997, provided guidelines for families that are evaluating intervention programs for children with autism.) Furthermore, a discussion of the general principles that constitute effective intervention is helpful. Dawson and Osterling (1997) furnished a review of the important elements in a child’s early inter-
vention programming, and Rogers (1998) surveyed the literature on empirically supported interventions for young children with autism.

Finally, although Josh's parents did not plan to have more children, their genetic risks were discussed. The team also presented the potential genetic risks of Josh's typically developing siblings having a child with an autism spectrum disorder.

**CONCLUSION**

Through this interdisciplinary evaluation, the team was able to offer Josh's parents a comprehensive and integrated picture of their son's current development and characteristics. By combining results across disciplines, his skills could be understood within the context of his overall development. For instance, his social, cognitive, and communication abilities were greatly influenced by his difficulty in sensory processing. This information would likely be useful to those working with Josh in the future. This case study provides the reader with information regarding assessment and diagnostic formulation of autism spectrum disorders in very young children as well. In addition, it illustrates one way of providing families with information that is coherent and useful while acknowledging what is not known in the field of early identification of and treatment for this disorder.

Given the importance and efficacy of early intervention for children with autism, an interdisciplinary team renders a crucial service. By providing families with information regarding the components of effective early intervention programs and by connecting families to appropriate services, the interdisciplinary team has an important role in changing the course of a child’s development for the better.
Case study 3

Excerpt from Virginia Walker Sperry’s,
Fragile Success: Ten Autistic Children, Childhood to Adulthood, Second Edition
Elizabeth Stanley: “I had to identify her emotions for her…”, Chap. 9, pp. 140–150

In early 1966, after several months of careful examination and study, the Yale Child Study Center told me that my 3-year-old child was different—“atypical” was the word they used. They went on to say that her atypicalness fell within the broad syndrome known as “autism.” I had never heard the term. They told me that she was born with built-in deficits, with no cause yet known. It was simply the way she was constitutionally constructed. This basic personality, along with its limitations, would be hers always. We had to accept and live with that.

I was terrified—at first by the cloud of mystery, and then by my gradual awakening to what it really meant, which was worse. Karen was this lovely baby we had so wanted, who was adored by all of us, including our four other children and relatives, who would never be normal. Since then there has been no significant disagreement among the different specialists we have consulted on Karen’s diagnosis and prognosis. All observed that Karen was not capable of imaginative play; she was withdrawn and unable to relate normally to other children. She was obsessive and perseverative. She was autistic.

By the time Karen was diagnosed, I had already known for some time that there was something wrong with her. When she was 2, our family lived for a year in a large city in the Orient. Before we left the United States, when getting the necessary immunizations at the pediatrician’s office, I hesitantly raised my hidden fears about Karen. Although none of her development was out of step with what I knew to be normal, she seemed slow: She wasn’t speaking; she felt limp; she lacked ordinary aggression (my other children at her age would have been all over the doctor’s office, climbing on chairs, or pulling papers from his desk). I said she seemed docile, too quiet. Was something wrong? The pediatrician mulled that over a bit, gazed thoughtfully at Karen, then reassured me: All children develop at different rates and over different spans of time; she was within normal bounds.

Our voyage abroad by freighter was a lively experience: 2 months at sea, visiting many ports in the Mediterranean, the Red Sea, the Middle East, and India. But I grew more troubled about Karen. Although I took two trunks full of toys, games, and books for the children, she touched none of them. She would not play with any of us, although she enjoyed it when we tossed her about and tickled her. The only amusement she found was splashing in the sink or bathtub in our room, and being carried about by the crewmen, who adored children.

After our arrival in the East, following the custom there, I employed an aide to take care of Karen. I also enrolled her in a small nursery group. But while all the others joined in play or listened to stories or sang songs, Karen sat on the fringe poking at the sand at her feet. She stood all alone, isolated.

After a while, we started to refer to Karen as “the quiet one.” We began to feel increasingly cut off, even rejected by her. At her age, after all, children look at you, hear you, laugh at and with you, and annoy you. Not Karen. I suspected that she needed medical help, and, so far from home, I became increasingly worried and at times nearly frantic.

When we returned in the fall of 1965 I immediately took Karen to the Yale Child Study Center to undergo professional scrutiny. While we awaited the outcome, friends and family tried to reassure us. But in those assurances I dimly sensed their need to protect me from some horrible truth they themselves had begun to fear. “After all,” they reasoned, “what do you expect? All of you dote on her and baby her to death. She doesn’t have to talk or do anything. Before long, she’ll get tired of that, and then you will see a big change. Einstein, you know, didn’t talk until he was 3.” We needed to believe that, so we did.

Meanwhile, we enrolled her in a nursery school, and out experience there prepared us for the reality the extensive testing would soon reveal. The teacher was completely baffled by Karen, so much so that she began to question her own competence. Finally, she insisted
Karen must be deaf. Worse, she worried that Karen didn’t like her and therefore would not respond to her as the others did. She felt hurt, a failure. I felt the same.

After 3 months of careful examination, observation, and consultation by the staff of the Child Development Clinic at the Yale Child Study Center, our doctor there told us the tragic truth. Karen was not deaf; she could hear a pin drop. This developmental specialist, her eyes full of dread her professionalism could not hide, said Karen suffered from autism, an affliction for which there was no known cure. Education was the only possible therapy.

The problem was that deficits in the formation of Karen’s personality were interfering with its normal development. Karen had no sense of her own identity, and she could not locate and use her own emotions. “You must get through to her, press yourself upon her, barrage her, make her hear you and respond. We are certain she can. You must reach her.” I could not miss the desperate insistence in the doctor’s voice.

The doctor went on. She said there was one other thing I had to do: I had to identify Karen’s emotions for her, and then interpret them for her and teach her what they were and the appropriate responses to them. I had no idea what this meant. For a long time Karen had seemed incapable of feeling or expressing any emotion. Now, the doctor was telling me that she really had emotions but could not understand what they were for or how to express them. Likewise, the doctor said that Karen had intelligence but could not use it. She could not relate normally to people or even to her physical surroundings.

As I listened, my mind whirled and my throat went dry. It all seemed unreal. I realized Karen’s responses and behavior were distressingly impersonal and sometimes bizarre. Now I was told I had to develop some way to force her to do what she apparently could not to do on her own—hear me, respond to me, deal with me, act like a human being. My God, what was she? Even dogs lick your hands, jump on you, invite you to play. Was she a sub-being of some sort? Could she even learn, respond naturally? What if she couldn’t? Would she become a “vegetable” and have to be “put away”? There are no words to describe those moments, the pain and terror I felt as I walked out of that office.

On arriving home, I opened the door. Full of apprehension, I went to the kitchen and put my purse on the table. Karen stood in the hallway. She acted as if I was not there. I looked at her. She did not look at me; I wasn’t “there.” “Karen,” I called softly, “come here.” She didn’t move. “Karen,” I said louder. She looked at the door. “Karen!” I shouted. She did not move. Tears of desperation filled my eyes. With both hands, I picked up a fat cookbook, lifted it over my head, and slammed it with all my might on the table, sending the salt and pepper shakers flying. “Karen!” I screamed. She turned and looked at me. She heard! I had reached her! She could respond! I was stunned. I knelt before her, gathered her in my arms, and sobbed. This was where I had to begin. What to do? Who would help? What would happen to her, to all of us, to me? Karen said, did nothing. So began the long trek to today.

Virginia Walker Sperry was director of the Elizabeth Ives School for Special Children from 1966 to 1972. From her years at the school, and through many to follow, she takes the reader into the lives of 10 of her students. From the beginning, where they experienced early diagnosis, medical and educational intervention, on into adulthood, Fragile Success shows us the determination not only of the students, but also the parents, teachers, and doctors giving their time and support. Her dedication, not only to the students education, but also their lives, shows the true spirit of Virginia Walker Sperry.
Karen and Our Family
At the very time in the life of a child when one expects the beginning of social interaction, Karen withdrew from us. She was rigid in body, fragile in bearing, vulnerable, and very frightened. All stimuli confused her unbearably and caused her either to recede or to engage in what we thought of as bizarre behavior. She seemed to be deaf. She found refuge in the bathroom, where she flushed the toilet endlessly; or she sat on the floor and rubbed the feet of a bronze statue we had brought with us from the East.

She immersed herself in things, in objects, in meaningless behavior. She was withdrawn, out of touch. Her gaze fixated at times on certain objects, such as the reflector pan of the stove or a crack in the table top. She screamed in terror when we blew up a balloon. She poked things repeatedly that caught her glance: a piece of jewelry on another person’s dress or a mole on the neck of a friend.

And the greatest problem of all for her future was her very poor reasoning ability and severe thought disorder. She had little to no ability for abstract thinking: She learned by rote, retaining what she held in her unusual memory. Pain, sorrow, anger—all these were beyond her. Such concepts as under, over, near, and far were like a foreign language to her, to be learned and routinely applied.

Her speaking as it emerged was often unrelated to anything and made little sense to others. She repeated a word or phrase over and over. As she learned speech, it was echolalic—automatically repeating what another person has just said.

She had no idea of spatial relationships or similarities, differences, or opposites. She seemed not to know that she had arms and legs, ears and eyes, or what they were for.

She insisted on objectifying anything (including living things) so she could touch it, handle it, mouth the words for it endlessly. She treated animate things as inanimate and inanimate things as animate. She turned people into things to be poked. She would pat and stroke a desk, or put a barn door hook-and-eye-latch in the baby buggy and push it about. Records were for spinning, not for listening. Dolls were for pulling apart. She even tried repeatedly to pull off her own head and stick it in the toilet.

She was tense and fearful. When frustrated, she would hit herself, pull up her skirt, or recede in a corner, thumb in her mouth. She dug at her fingers until they bled.

In all, this one fact stood out: She could not relate to people as people. Locked inside this vacant isolation, Karen was virtually unreacha by anyone, beginning with her family, and of course I feared she could eventually back out of human contact entirely and into a world none of us would be able to enter. The prospects were horrifying.

Worst of all, she did not know us, who we were, or that she was our baby girl and our other children’s sister. She had no idea who she was, that she was a girl, a daughter, a sister, a playmate, a person. She did not play with her siblings or delight them, as babies always do. This was, I felt, a loss for my other children beyond speaking. Likewise, she did not know me as her mother: I had no special meaning for her. She could easily walk off with the next stranger on the street, and sometimes did. She didn’t know the difference. It was years before she knew who her parents and siblings were, and where and what home and family were. All of us felt rejected.

I was nearly crushed. If you can penetrate the heart of a mother and perceive what really feeds her inner being and makes her enjoy being a mother, it surely includes the wonder and excitement and ineffable satisfaction of the response of her child to her. But
this could not be; it was impossible to mother Karen. I was screened out. I had no cue from her to go on; I was at a dead end. I could only do things for her or to her and see that she was fed and clothed. The distance between such care and mothering is vast.

In those early days, I could think that I, in some way unknown to me, had caused her problem. I am sure every mother of an autistic person suffers this torture. But in time I came to understand that the cause stemmed from Karen’s own constitutional makeup and her consequent development. Slowly, I began to realize that these symptoms of autism were not rejection of me or of us but rather her escape from impossible-to-endure confusion and inner turmoil over which she had no other control. The symptoms were, in fact, her very means of survival. We would have to find out how to get around or through them to help her take control and release the real person within.

**Parents and Family At Risk**

The difficulties an autistic person brings to a family are enormous, especially for the mother. You go on being a wife, a mother to the others, a neighbor, a friend, a relative—taking up those relationships while you try not to allow the autistic person to absorb you to the detriment of your life. You try to avoid feeling guilty, for if the mother feels guilty the problem is only compounded. And no matter how much you give, the autistic person still needs more.

Siblings often feel abandoned by a mother given over to the care of an autistic child. They feel neglected, or worse, they feel guilty that their demands might increase their mother’s already too-great burden. It is easy for them to fear that adding to their mother’s worries might push her over the brink. They do not feel free to express what might be ordinary discontent about their situation.

In such circumstances, how can a normal adolescent work through the ambivalence and complexities of growing up? This was especially true in the 1960s and 1970s when Karen was young, which was a time of great social upheaval and disenchantment of the young with the establishment of which we, the parents, were of course a part. So, Karen’s siblings became mute and tended to go their own ways, missing sorely what they needed from their parents.

Their apparent alienation made things difficult for me, too. Because they were normal, I expected them to do a lot better for themselves and to help with Karen. It was very easy for me to feel everyone left me with the burden, isolated and alone. Such feelings on my part only increased their worries and drove them further away. And the problem worsened.

My husband was usually not directly involved in the day-to-day care of Karen. Some men, my husband included, find the pain of having and caring for an autistic child simply too great to live with intimately, and still carry on their work. I now see that it might have destroyed him, as he feared it would me. He had to protect himself, and this meant distancing himself as he could from the everpresent trouble.

My husband did help where he could—with the transportation of the children to their social and school obligations and of Karen to her numerous appointments and programs. He did keep steadily at his job and its demands and supported the family. He sometimes made it possible for us to get away alone where we could restore ourselves and be refreshed. He involved himself in every decision. Although relating to Karen baffled him, he cared about her and in one way or another built a strong relationship with her that today serves her well. Now, he delights in her accomplishments, growth, and presence and participates in the new challenges we face with her. He often picks her up at her residence to take her shopping and to dinner at a restaurant, a treat she always accepts with much pleasure.
Once, when speaking to a parent group, my husband said he often felt as though we were in a lifeboat. Karen had fallen overboard and I was about to jump in after her, even though I didn’t know how to swim. His job, he said, was to steady the boat and keep me from jumping. (It is the mother, after all, who jumps. How could I possibly sit there and watch my child drown?) He once told me flatly, “If it is a choice between you and Karen, I choose you.” Well, in a way I was reassured and in a way threatened. What did choose mean? Giving up Karen? I had to keep Karen and myself stable so that such a choice would not be necessary. So I kept quiet.

And in that is the essence of the deepest isolation of all the mother of an autistic child feels. I couldn’t imagine giving up on Karen, yet, I couldn’t express to my husband or children what I was feeling, for I knew it would hurt them and drive them away from me. So, I felt increasingly alone, trapped. I have since learned that most mothers of autistic children do.

Karen’s Education

At the time Karen was diagnosed, there was only one small school in our area that was trying to deal with autistic children: the Elizabeth Ives School for Special Children. The doctors at the Yale Child Study Center suggested we enroll her there, which we did.

In all, Karen has gone to five schools, some of them new and experimental. From her earliest years, schooling for Karen followed a consistent pattern. She had a succession of wonderful teachers who were committed deeply to her, and who believed she could be made to hear them and to learn. Many of them believed firmly that she had a concealed intelligence and emotional sensitivity, which they sought desperately to reach and to activate. These forceful and persistent people provided Karen with the necessary structure and support for her to begin to participate in life, to respond, to speak, to know herself and to let herself be known.

Fundamental to the method used was the drive to get a reaction from her. They, the teachers and we, the parents, demanded a relationship with Karen, and in demanding it from her, we began to get it. She had to look into her eyes, hear us and respond to us. We insisted. It was a gut job, not a paper exercise. It made us sweat. Teaching her reading, writing, and arithmetic at that stage seemed less relevant than nurturing a relationship in which we helped her to identify pain, anger, affection, humor, hurt, and grief and express them appropriately.

Karen’s education and training went on at school and at home. Her disabilities forced us to teach her survival skills—what she had to know and to adapt to a world she could not understand. We demanded, so far as we could, acceptable behavior. She gradually learned to look after her personal care, to take her own shower, to tidy her room, to understand and manage menstruation, to dress nicely, to greet people. Each acquired skill prepared her to learn the next and to grow in the art of being human.

There was an additional, extremely potent, force at work for Karen. It lay within her. She wanted to relate to us. She wanted to learn and be independent. All the while she was growing up, she had not been really so out of touch as her withdrawal had made it seem. As time and schooling went on, the layers of resistance and withdrawal peeled away, and we learned that she wanted to laugh, to sing, to dance, to be with us, to draw, to pile up blocks and knock them down, to swim, to walk, to listen to records, to watch the birds, to learn the names of flowers and trees, to go places and do things.

Speech came awkwardly. At first it was hollow and regimented, then slowly her own speech emerged, clear and crisp. She learned to assert herself and defend her possessions and her privacy. She wanted friends. As she learned to express her pleasure in people she
began to elicit the affection of those with whom she came in contact. Her smile took over and the pleasant person she is emerged. For all her stilted speech, she won interest and affection, and more and more people made their adjustments to her difficulties and peculiarities and related to her. They not only made her happy, they helped her learn to live her life. It all came with glacial slowness. It was only after long spans of time had passed between what she had been and what she was becoming that we knew progress was being made.

As Karen learned to relate to people, her ability to learn the fundamentals of the simplest survival skills and elementary academics improved, and her inner disturbance began to give way bit by bit to inner calm. She began to take control of herself and emerge as a person. She learned more and more of the social skills that are essential to normal living. In time, we found that she could not progress very much academically. We had to learn when to stop our pressure.

Over the years, Karen has been examined, evaluated, and treated by competent psychologists, psychiatrists, social workers, and educators. We have employed helpers and teachers privately. Connecticut during these years was in the forefront in taking governmental responsibility for the education of all children, even those with impairments as serious as Karen’s. So, happily for us, the state bore part of the horrendous cost of educating and treating her.

**Adulthood and Supported Independence**

After many years living at home and in group homes, Karen now lives with two other women with developmental disabilities in a semi-supervised apartment in a nearby city. The state DMR provides 20 hours a week of staff supervision for this semi-independent living program. In Karen’s case, the staff person is a capable adult who helps Karen and her apartment mates organize their home, provide food and necessary supplies, schedule medical visits and plan and take part in social and recreational activities. A vocational counselor helps locate jobs in the community. Thus, Karen is more and more thrown on her own resources, and often she is able to develop her own skills to meet her emerging needs. However, she still needs much help—the most difficult sort to provide—in taking responsibility for herself, and in planning and executing the means of satisfying her needs.

Given her basic impairments and the long-term care by staff members, her progress toward increasing independence is slow. She still has no concept of numbers or money, and quite confidently empties her wallet in front of store clerks, who are then expected to take only what is needed. So far, her trust has not been betrayed. But this illustrates how much we who parent such autistic people, must trust the fragile and often dangerous society into which we launch our children, hoping not only that they can manage, but that others will give them a hand and not harm them. With Karen, our trust has been rewarded, but I can never give up fearing for her. Even crossing the street is a major hazard because with cars coming, she will start across the street, all the while assuring me she has “looked” and, having looked, is free to go. Looking, assessing what she sees, and transferring that into appropriate action is still a very complicated process. For Karen learns not so much by instruction as by immersion. I was never one to throw a child into a swimming pool and say “swim,” but now I do it all the time with Karen. There is no alternative if she is to be able to live in the world as it is. For that goal I am willing to risk her life, to take the chance that she will make it. This is not a testimony to rashness but to my desperation and her need.

Any program in semi-independent living for an autistic person is full of risk and uncertainty. Karen is frequently on her own, busying herself at her home, traveling over the one
bus route she knows to a mall, which, I have learned, is like a village green, opening to take her to the world of shops and the possibility of meeting a friend. She takes up formerly impossible tasks such as getting herself bathed, dressed, and outside ready to meet a van that takes her to work each morning. Even managing not to lose her house key and actually being able to put it in the lock, turn it, and open the door without a helper in sight is a major step ahead, and a building block upon which she develops more confidence and other skills. The two people who share her apartment are good friends, and they all help each other in many ways. Their apartment has become a kind of informal gathering place for others in nearby semi-independent programs. They celebrate birthdays, go out together to special events, send out for pizza, and watch TV and videos. They hang out.

Efforts are made to help Karen and her friends explore and use resources in the community, such as churches, YMCAs, health centers, libraries, movies, concerts, and eating places. Like most adults, they move about in a circle they come to know and can handle on their own. Up until 2 years ago, Karen lived in group homes under 24-hour supervision and had never been left alone 1 day in her life, even at home. Until she was 30, her entire life was programmed, supervised, and carried out under close staff supervision. She could not use a telephone or take a bus or taxi. She could not prepare her own meals, organize her daily life, plan and carry out wanted to do, or pay her own bills.

Now she can use the telephone and most of the time, dial it correctly. She helps clean the house and does her own laundry. She can open canned food and microwave TV dinners. She is often alone in her apartment, even for an entire weekend, sometimes preferring that to coming home to her parents. She can take the bus on a limited basis; she can call a taxi when needed. She can walk alone down her street a couple of blocks to a variety store to buy milk and bread. On Sunday she takes a taxi to church (having telephoned the previous day to arrange to be picked up). She goes to work each morning, still in a semi-supervised job, but increasingly on her own.

Karen still has major deficiencies that would confound any normal person. She cannot handle the telephone reliably: She will frequently pick up the receiver and not say anything, expecting it to talk to her. She does not get messages straight. She cannot read or write beyond primitive levels. She gets lost if a well-defined route is interrupted. She does not approach others to ask for help or directions. Her recreational outlets are few, her friends limited to those at least as limited and impaired as she is. When frustrated, she may “act out” by screaming and breaking things, not yet knowing how to express her feelings appropriately and work out her frustrations in a more useful way.

As inadequate as it sometimes seems, I know Karen’s life depends on the resources the state can give her in living independently and learning to manage her own life. Although I cannot see very far down the road, I am sure she can grow more and more into a life she can manage on her own. Increasingly, she will become independent, a goal once so far beyond the realm of possibility. Never in my wildest dreams did I imagine it could happen. Well, it hasn’t yet, and every week is fraught with problems. But, we muddle through, and in the end, I am sure Karen will survive as her own person in her own life.
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The mind is mysterious, especially when it comes to autism. A person with autism might lack the ability to put together a complete sentence, but at the same time exhibit gifts that outreach the grasp of most people. Such autistic savants commonly excel in art, music, and math. Often, their skills are not cultivated through instruction but are self-taught.

Meet 16-year-old Jonathan Lerman from Vestal, New York. Despite his limited communication skills, Jonathan is able to convey through art a world that stretches far beyond the confines of language. Although autistic savants have gained a lot of attention through films like Rain Man, only an estimated 10% of people with autism display savant abilities. Nevertheless, according to Stephen M. Edelson, Ph.D. at the Center for the Study of Autism, people with autism tend to be 10 times more likely to possess special skills than people with other developmental disabilities.

The Artist Within
At the age of 10, Jonathan suddenly began drawing. He had just lost his grandfather with whom he shared a special bond. Upset over his grandfather’s death, Jonathan would often ask Caren when he could visit him in heaven. Equally fond of Jonathan, his grandfather used to tell Caren that Jonathan had promise and that one day he would surprise her. Apparently, he was right.

In 1997, Caren received a call from Jonathan’s helper at the local Jewish Community Center. “You’ve got to come over to see what he’s doing,” she said. With trepidation, Caren replied, “What, holding the other kids hostage?” “No, he’s drawing,” came the answer. Jonathan was not only drawing, he was doing so with the ease, swiftness, and confidence of a master, turning out 5 and 10 drawings in one sitting. Even more remarkable, he was drawing faces complete with sorrow, happiness, and the vast array of emotions that make up the human psyche. It is something that no other recognized artist with autism has ever done before. “He totally broke the mold,” explains Caren.

Jonathan’s work has met more than his mother’s standards. Art critics and professors have also proclaimed its excellence. In a New York Times article, John Thomson, chairman of the art department at New York State University at Binghamton said Jonathan’s drawings “would not be out of place in my classroom.” Thomson added that it was “really exceptional, characterized by an amazing lack of stereotypes common to drawings of all age levels.”

Living with Success and Autism
Jonathan wasn’t always tuned into people and faces. In an unpublished memoir, “The Solitary Heart,” Caren described Jonathan’s anti-social behavior as a child. “When our friends would come to our house, he would find a wall, lie along its side and stare at it for hours.” Similarly, at the beach, Jonathan would step on sunbathers with the same gusto that he would step on sand.

Now, Jonathan isn’t just noticing people, people are noticing him. K.S. art gallery in New York City has featured
his work in several solo shows. Ranging in price from $500 to $1800, about 70 of his drawings have sold so far. Charcoal is Jonathan’s preferred medium but he occasionally uses pastels. Over the last couple of years, Jonathan has ventured into working with clay. Caren confesses, “We’ve kept all the sculptures so far. They’re just so amazing, but we plan to share them and market them in the future.”

Caren is thrilled about her son’s talent and success but she admits that raising Jonathan wasn’t easy. Still, she says, “[Raising a child with autism] has taught me compassion, patience, and the value of unconditional love.” To view samples of Jonathan’s artwork or to purchase Jonathan Lerman: The Drawings of a Boy with Autism, visit www.ksartonline.com.