The Inner History of Devices

edited and with an introductory essay by Sherry Turkle
### Contents

viii Acknowledgments

**READING THE INNER HISTORY OF DEVICES**

2 Inner History | *Sherry Turkle*

**THROUGH MEMOIR**

32 The Prosthetic Eye | *Alicia Kestrell Verlager*
41 Cell Phones | *E. Cabell Hankinson Gathman*
49 The Patterning Table | *Nicholas A. Knouf*
55 Television | *Orit Kuritsky-Fox*

**THROUGH CLINICAL PRACTICE**

64 The World Wide Web | *John Hamilton*
77 Computer Games | *Marsha H. Levy-Warren*
86 Cyberplaces | *Kimberlyn Leary*

**THROUGH FIELDWORK**

98 The Internal Cardiac Defibrillator | *Anne Pollock*
112 The Visible Human | *Rachel Prentice*
125 Slashdot.org | *Anita Say Chan*
138 The Dialysis Machine | *Aslıhan Sanal*
153 Video Poker | *Natasha Schüll*

172 Notes

198 Index
transferred with the rest of my telephone contacts. My dead relationship's ring tone is not.

And it's not just his ring tone that's gone. He belongs to another artifact. It sometimes feels as though the year of my life I spent in that ill-advised love affair was poured into the now-obsolete Audiovox. Like a gift that still possesses something of the soul of the giver, the phone itself had come to be haunted by the voice that spoke so often through it. I gave back everything he had given me, but I couldn't get him out of my phone. I used to get tattooed when love went wrong, remaking my body into one that had never known the touch of my former partner. Now I have a new phone into which I have never spoken too soon or not wisely enough. Now I have a phone from which I have never failed to hear the words I wanted. My new RAZR is a part of a me that is freshly born, unscarred.

And I carry it everywhere. I frequently speak on it as I move through public places—I find that it provides insulation against panhandling and unwelcome advances. It's difficult, after all, to engage someone in conversation when they are already talking to somebody else. If I can't get a signal, I don't go as far as my friends who will pretend to talk to a dead phone; I just play Tetris instead. The blonde girl with her pearlescent pink _keitai will never walk the streets of Tokyo again, but now there's a pink-haired woman with a pink zebra-skin RAZR that trills "Heartbreak Beat" for default calls in Boston.

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**The Patterning Table**

Nicholas A. Knouf

Our basement walls were covered with charts, schedules, and sets of instructions. In the southeast corner stood the patterning table. Made when I was eight years old by a friend of the family who lived in a nearby town, the patterning table came up to my chest. Supported on its four corners by thick wooden legs attached with massive bolts, the highly varnished table was strong enough for a fifty-to-eighty pound person to be moved around on its flat surface. That surface was padded with Naugahyde, which in turn was covered by sheepskin that fitted snugly around the edges.

At this table, every Monday through Saturday, three people (five, if we were training new volunteers) surrounded my sister, Robin. We gently took hold of her fragile arms, legs, and head. With a regular rhythm we moved her extremities in the motions of a crawl: one, turn the head to the left, bring the left arm away from the body and next to the head, bend the left leg next to the torso, and vice versa for the right side; two, keep the head where it was, pull the right arm back next to the body, extend the left leg, and vice versa for the right side; three, repeat as one, but switching left for right; four, repeat as two, but switching left for right. One, two, three, four, the count continued over the course of five minutes, and the patterning session was done, for this hour. The ding of a kitchen timer told us it was time to move on. Next hour we repeated it, on
the same table with the same sheepskin cover, with different volunteers. All day this continued, according to the schedules, instructions, and charts that my mom wrote in thick strokes on butcher paper and taped to our recently sheetrocked basement walls. Patterning was combined with breathing exercises, using a mask designed to increase the blood flow to Robin’s brain.

When I was in first grade Robin was diagnosed with Rett Syndrome, a rare neurological disorder that may include devastating mental and physical retardation. It afflicts only girls, most of whom are not expected to live past their eighteenth birthday. The girls often make rocking motions while sitting, wringing their hands. Many are able to develop some basic level of functioning, such as simple mobility or being able to feed themselves. Robin, however, could do none of these things; she was utterly dependent on us. Even so, there were still the smiles, tears, and frowns of any young girl, and in her eyes you could see thoughts she could not speak.

Robin’s disease led our family to take on a major commitment: a full-time, in-home physical therapy program. Our location, rural Iowa, meant we had to develop everything ourselves. To build the equipment, to schedule the people who came in on a regular basis, week after week, we found volunteers, some local, some from afar.

The patterning table took up a good part of our basement, but at a certain point during Robin’s therapy we set up another apparatus for her, a jungle-gym-like contraption made by the same volunteer who built the patterning table. The new contraption reached from the floor to our seven-foot ceiling. The height of the jungle-gym bars was adjustable. At certain times the overhead bars were low enough so that Robin could walk her hands along them. When she was stronger, the bars were raised and Robin had to grab onto freely swinging handholds of thick nylon rope that hung from the bars; she wore specially designed Velcro footies that stuck to our short-pile institutional carpeting. The resistance from the Velcro was meant to make it harder for her to lift her foot, while the swinging rope was meant to improve her sense of balance. It was hoped that this would strengthen Robin’s legs and upper body in preparation for walking on her own.

Our work at the patterning table was based on the Doman—Delacato method. It laid out a series of therapies designed to help brain-damaged children toward better functioning. The method is based on an evolutionary metaphor: the individual develops in their lifetime just as the human evolved over generations. That is, development begins at the fish and reptilian stage (crawling) and moves through to mammals and primates (creeping, with the stomach off the ground). Doman-Delacato reasons that if you “pattern” a brain-damaged child with the motions that are involved in each of these stages, you will unlock the later stages of development. So the repetitive motions on the patterning table were supposed to teach Robin’s muscle memory to crawl. Yet she never was able to crawl on her own: she learned only to walk a step or two, lightly aided.

The Doman-Delacato method has not found empirical support, and the American Academy of Pediatrics has issued a number of warnings about the technique over the years. As a child I never questioned our program. To me, it seemed natural that if we moved Robin’s arms and legs in a certain manner, we eventually would “train” her brain to move body parts on its own. And of course my parents, like so many others, seized on the program’s marketing and the bits of anecdotal evidence that suggested it worked. The Doman-Delacato method provided hope, but I think back on it now with sadness, regret, anger, and resignation.

Robin began to experience seizures when I was in fifth grade, not unusual for Rett girls who often regress.
from their developmental plateau. I remember the huge stuffed rabbit that lay on the table the day Robin was rushed to the hospital because of her first, unexpected seizure. The seizures made it impossible to continue with the treatments on the patterning table. Yet the patterning table remained in the basement. We removed the sheepskin, and then the Naugahyde quilting, and used the table as any other flat surface, a place to store papers, books, and mail. Gradually the schedules and charts on the butcher-block paper began to come off our walls, but the patterning table remained, a reminder of what we had done, of what we had tried to do.

Even as a young child, I read the few journal articles our family had about Rett Syndrome. They were over my head, but I persisted in my efforts to figure out what was going on with my sister. I launched into my parents' popular science book on genetics. A year or two before I took any proper biology class, I was discovering genotypes, phenotypes, and karyotypes, the genetic bases of diseases such as sickle-cell anemia and Parkinson's. At the time, the Human Genome Project was in its nascent stages and the hope for "miracle" cures was strong. With the insufficient knowledge I had, and bolstered by the arguments in the genetics book, I believed that finding the gene or genes that "caused" her disease would lead immediately to gene therapy that would "reverse" Robin's genetic malfunction. Of course you had to have a way to get the genes into the cells, so I eagerly read about techniques to force the existing chromatin to take up new genetic material—retroviruses and exotic "electro gene therapy." Then came the problem of how to reverse the damage in existing cells. Neuron regrowth in the brain is meager at best; there, repairing genes would not be enough.

As a middle school student growing up with my sister, I thought these problems surmountable. And I determined that all of this would be my doctoral work:

I would find the gene that caused Rett Syndrome and discover the necessary therapies to cure the disease. It was merely a matter of time. The summer before high school, I enrolled in a summer program in molecular biology at the state university, a first chance to work with the tools I had read about: restriction enzymes, plasmids, and ethidium bromide stains. I was in the throes of passion: I pored over books I could not understand; I persuaded other students to work unattended in laboratories full of dangerous reagents. I was in a hurry.

But Robin died in the fall of 1993, just short of her ninth birthday. I sunk into a world of grey and black, my schoolwork the only thing that kept me going. I remember the fleeting thought that if I could complete my work faster, I could start college early and be on my way to finding the cure for other Rett girls.

I went to Caltech as an undergraduate to study molecular biology with one sole objective in mind: to find a cure for Robin's disease. In high school I had performed in a chamber music group and had notions about attending a conservatory. I devoured literature and thought about being an English major and studying philosophy. But now, stronger than all of these were my memories of Robin and the patterning table. I was driven to discover the gene for Rett syndrome. I saw it as an achievable goal.

Then, on a gray California winter day during my sophomore year, I read that a group of researchers had discovered the gene responsible for Rett Syndrome. I posted the article outside my room. I told all of my friends. I wanted to be among the people in the article. But there was more work for me: turning the genetic discovery into a treatment for Rett patients. Soon after, experience as a research assistant convinced me that I
The patterning table, long unused even during Robin's life, was finally removed from our basement, probably passed on to another family using the Doman-Delacat method. Its absence left a space. I had passed the patterning table and its volunteers in the early morning; I had looked toward it when I came home from school; I had walked to it when it was my turn to be part of the group that moved Robin's head, legs, and arms. Without the steady presence of the table, where would we turn? Where would our efforts be channeled? Without Robin's influence, what would be our purpose? The table was more than a focus for our thoughts; it anchored our love for Robin and the energy we put into giving her a future. Like the tables in traditional Midwestern churches or cafes, the volunteers who assembled chatted and gossiped and spoke about their lives. It brought Robin into a community. When Robin was happy and obedient, her attendants gave her praise; when she was cranky and ornery, they understood, but lightly scolded. The patterning table made life coherent, all of a piece. With Robin, the volunteers, the discredited therapy method, and the patterning table, we had tried to awaken cognition with care, with the soft sheepskin, the men and women gathered around.

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The first time I watched television I was four; my mother and I went to visit the downstairs neighbors. On the screen of their newly acquired TV set, people were stepping into mirrors and reemerging on the other sides, into new rooms with different mirrors.

They tarried there long enough to explore, and then crossed back to their starting points. I held my breath every time these people were at a threshold, imagining my own skin submerged in the mirror's thick grey mercury-like substance. When the people safely passed to the other side, I inhaled deeply, as if emerging from a dive in deep water. In the years that followed, I would do that every time I watched one of this program's mirror-crossing segments, a program I later learned was called Vision On and produced by the BBC. I have no memory of what the neighbor's television looked like. I can recall it only as a blurred shape, reduced to another mirror frame whose purpose was to be passed through.

But I do remember a twist in that first episode: one of the characters—a tall, bearded man—walked into an elongated mirror in his living room, but to my horror, when he crossed back, he found himself in a different room. He continued through another mirror into another room, and then another and another. The chain of mirror crossings ended outside, on a manicured lawn. The scene now dissolved to a slapstick sequence: someone (or something) was chasing the bearded man, and