Developing a Program for Training Therapists to Conduct MDMA-Assisted Therapy

MAPS is assisting researchers to design, fund, and conduct the clinical trials necessary to develop MDMA-assisted psychotherapy into a recognized treatment for posttraumatic stress disorder (PTSD) and/or clinically diagnosed anxiety secondary to life-threatening illness. Currently, MAPS is sponsoring several Food and Drug Administration (FDA) Phase 2 studies of MDMA-assisted psychotherapy in people with PTSD and anxiety secondary to a life-threatening illness. Studies of MDMA-assisted psychotherapy in people with PTSD are taking place in the United States and Switzerland, with a study in Israel to be initiated this year, and a study of MDMA-assisted psychotherapy in people with anxiety associated with an illness is under way in the United States. If findings from these small pilot studies continue to be promising and indicate that MDMA-assisted therapy is safe and possibly efficacious, then MAPS must develop and conduct two large-scale multi-site Phase 3 studies if we wish to apply to FDA for approval and recognition of this treatment. Each of these studies will involve approximately eight teams of psychotherapists, and approximately 280 participants. Each therapeutic team will treat as many as 18 participants per year for two years. One Phase 3 study will be conducted throughout the United States and the other throughout Europe and Israel.

Such large studies will require training many mental health professionals to conduct MDMA-assisted therapy. This is a situation for which we know of no precedent.

The MAPS team is in the process of developing such a training program, to be led by Michael Mithoefer, principal investigator for the first study of MDMA-assisted therapy for people with PTSD. The training program will provide therapists interested in conducting MAPS-sponsored MDMA-assisted psychotherapy research with the experience, education, and information needed to perform this therapy.

Currently, under the training program we envision, we will enroll pairs of therapists in male-female co-therapist teams, matching MAPS’ treatment method for MDMA-assisted therapy, which requires a pair of therapists and strongly encourages the pair to consist of a man and a woman. Right now, the most likely design will have one or two teams of therapists-in-training visit Charleston, SC, on two separate occasions to meet with Michael and Annie for a four or five day training event. During this time, the trainees will each experience one session of MDMA-assisted therapy as the designated patient, with their partner as the therapist, and one session as the designated therapist, with the partner in the patient’s role.

The second visit to Charleston will be scheduled at least three months after the first so that the trainees have the time and opportunity to observe and learn not just about the MDMA experience itself but also how to integrate it into their daily lives.

The training program might involve bringing as many as 10 teams of therapists together for a week-long educational seminar about the use of MDMA-assisted psychotherapy for PTSD after they have all completed two training sessions, allowing for greater learning and sharing of experi-

Ilsa Jerome, PhD, (digital self-portrait), ilsa@maps.org and Rick Doblin, PhD

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ON MARCH 19, 2007, during a long phone call with the president of the Ethics Committee (EC) (the Swiss equivalent to an institutional review board), which is responsible for the approval of my proposed MAPS-sponsored study, she announced that the committee had granted conditional approval.

So we have reached a milestone. I am convinced that the approval by the EC is the most difficult step in the entire approval process. Ethical decisions are judgments. Four years ago I was a member of a group of researchers from the Swiss Medical Society for Psycholytic Therapy (SaePT) that had a frustrating experience with a psilocybin/depression project that was not allowed to proceed following rejection by the EC.

During the recent EC meeting, although the committee was critical and posed detailed questions, in general they were not overcome by prejudice. In the end, the committee was convinced that the potential benefits of LSD-assisted therapy outweigh the risks.

Now, how to continue? First, before receiving unconditional approval, I have to wait for the written report of the EC and fulfill their requirements. Then, I will submit my papers to Swissmedic (Swiss Food and Drug Administration equivalent) and finally to the BAG (Swiss Drug Enforcement Administration equivalent). After all three of these groups grant approval, I will have full regulatory approval for the study.

I am happy and relieved to have reached this step. I’d like to thank everybody who supported me until now. As a researcher in private practice I depend on a network of people who are able to support this work. I’d like to thank Rick Doblin for his enthusiasm and financial support; Ilsa Jerome for busy and patient support in scientific literature research; Valerie Mojeiko, Amy Emerson, and Josh Sonstroem for their methodological support; John Halpern and Matt Bagott for permission to use their protocols; Rudolf Brenneisen for his support in accessing and handling the LSD; and, finally, I thank Albert Hofmann for the opportunity to consult with him about this study.

This was a big step in the right direction, although it’s still a long climb to the mountain top. I’ll keep you up to date! •