Reston, Virginia, March 30, 2005—Jacqueline D. Griffiths, M.D. of NewView Laser Eye, Inc. in Reston, Virginia is pleased to announce today that the Moria Epi-K™, its disposable epikeratome for Epi-LASIK, has received FDA approval in the United States.

The Epi-K™ is utilized to mechanically separate the epithelium from Bowman’s membrane. The epithelial flap is then folded back prior to laser ablation, and subsequently returned to its original position. The procedure preserves the structural integrity of the stroma and is expected to minimize discomfort, shorten the length of visual recovery, and reduce the incidence of haze associated with other surface ablation procedures, such as PRK and LASEK.

In rigorous clinical trials at 13 sites in 9 countries on over 500 eyes, the Epi-K™ achieved excellent results. All clinical investigators reported that the device produced very high quality epithelial flaps and that postoperative pain and visual recovery compared favorably with other surface ablation procedures. Dr. Barrie Soloway, principal investigator at one of the US sites, noted that 88% of patients indicated they could return to work within three days following surgery.

NewView Laser Eye, Inc. is the first center in the area to offer this technology. Dr. Griffiths is excited about the ability to offer patients the latest advances in laser vision correction. Please call us at 703-834-9777 or toll free at 1-800-294-1001 for a consultation or with questions. Or, visit our web site at www.newviewlaser.com.