BOTULINUM TOXIN TYPE A
CONSENT FORM

Patient Name: ________________________  Date: __________________

To the patient: Being fully informed about your condition and treatment will help you make the decision whether or not to undergo Botulinum Toxin Type A treatment. This disclosure is not meant to alarm you; it is simply an effort to better inform you so that you may give or withhold your consent for this treatment.

I have requested that ____________________ attempt to improve my facial lines with Botulinum Toxin Type A. These injections have been used for more than a decade to improve spasm of the muscles around the eye, to correct double vision due to muscle imbalance as well as numerous other neurological uses. Botulinum Toxin Type A is now approved by the FDA to improve the appearance of the vertical lines between the brows. A few tiny injections of Botulinum Toxin Type A relax overactive muscles and soften those vertical lines. Injections in other areas to improve appearance of facial lines have been reported in the literature, but the FDA has not approved those uses. The results of Botulinum Toxin Type A are usually dramatic, although the practice of medicine is not an exact science and no guarantees can be or have been made concerning expected results.

_________Patient Initials

The Botulinum Toxin Type A solution is injected with a tiny needle into the muscle; you should see the benefits develop over the next two to seven days. A decreased appearance of frowning or creasing of other lines will be the result of this treatment.

_________Patient Initials

The most common side effects are headache, respiratory infection, flu syndrome, temporary eyelid droop, and nausea. Botulinum Toxin Type A should not be used if there is an infection at injection site. Additionally, slight temporary bruising may occur at the injection site. I have been advised of the risks involved in such treatment, the expected benefits of such treatment, and alternative treatments, including no treatment at all.

_________Patient Initials

I understand that the results are temporary and several sessions may be needed for optimal results.

_________Patient Initials

I acknowledge that I have received literature with full disclosure regarding any and all possible side effects or complications from use of Botulinum Toxin Type A.

_________Patient Initials

I agree that this constitutes full disclosure, and that it supersedes any previous verbal or written disclosures. I certify that I have read, and fully understand, the above paragraphs, and that I have had sufficient opportunity for discussion and to ask questions. I consent to this Botulinum Toxin Type A treatment today and for all subsequent treatments.

Patient’s Signature: ______________________________ Date: ________________

Witness Signature: ______________________________ Date: ________________