National Prion Disease Pathology Surveillance Center
Testing and Reporting Policies

As a part of our surveillance efforts for CJD, the National Prion Disease Pathology Surveillance Center (NPDPSC) conducts four different tests on the biopsy and autopsy samples we receive:

- **Western blot:** This test demonstrates the presence of the abnormal prion protein, which is believed to cause CJD and other prion diseases. If the abnormal protein is present, the case is positive. The Western blot is the most sensitive test for prion disease. This test is performed on frozen tissue.
- **Immunohistochemistry (IHC)/Histology:** In these tests, the neuropathologist examines slides of specially prepared brain tissue to see where the abnormal prion protein appears in order to help determine the type of prion disease. Different types of CJD have different distribution patterns of the abnormal protein. These tests are performed on fixed tissue.
- **Genetic analysis:** This test determines if the patient has a genetic mutation, and therefore a familial prion disease. The genetic analysis can only determine if a case is familial (which occurs in about 10% of positive cases); in all other forms of prion disease such as sporadic, iatrogenic, or variant CJD, the genetic analysis may help to identify the specific type. This test is performed on frozen tissue or blood. If we receive sufficient amounts of frozen tissue, blood is not required.

All four of these tests must be performed in order to provide a full diagnosis. We perform all of them providing that the appropriate samples are provided. If one of the samples is not available, we cannot perform all of the tests mentioned above and cannot provide a full diagnosis.

Although we perform all of the above tests for our important research efforts on prion disease, we realize that some families may not want all of the information these tests provide. In particular, some families do not want to receive genetic information. Genetic mutations not only affect the patient, but other blood relatives who could also have the mutation. It is important to discuss the psychological implications, confidentiality and insurance concerns with them to determine if they wish to receive this information.

In order to ensure that the family receives only the information they would like, we are asking clinicians to consult with families to determine if they would like to receive a full or partial diagnosis. Please indicate their choice below and fax it to us at 216-368-4090. The NPDPSC will not release genetic information until this form is returned with physician signature.

For blood only cases where the family wishes to receive the genetic information, please check the “full diagnosis” box to release the genetic analysis.

For questions, please contact us at 216-368-0587 or cjd@case.edu.

- Please check the appropriate box listed below:
  - [ ] Please send only a partial diagnosis, including the Western blot (if frozen tissue is available) and IHC/Histology (if fixed tissue is available), without the genetic analysis. The partial diagnosis will only indicate if the case is positive or negative.
  - [ ] Please send the full diagnosis, including the genetic analysis (only available if blood/frozen tissue is submitted). The full diagnosis will indicate if the case is positive or negative and provide the type (sporadic and the subtype of sporadic, familial, or variant) of prion disease if the case is positive.

Patient Name: ___________________________ Date: ___________________________

Physician Name (print): ___________________________ Signature: ___________________________

Physician Phone: ___________________________ Physician Fax: ___________________________