Product Information

Monoclonal Anti-Factor IX antibody
produced in mouse, clone HIX-5
purified from hybridoma cell culture

Catalog Number F1020

Product Description
Monoclonal Anti-Human Factor IX (mouse IgG1 isotype) is derived from the HIX-5 hybridoma produced by the fusion of mouse Sp2/0-Ag14 myeloma cells and splenocytes from BALB/c mice immunized with factor IX purified from human plasma. The isotype is determined by a double diffusion immunoassay using Mouse Monoclonal Antibody Isotyping Reagents (Catalog Number ISO2).

Monoclonal Anti-Factor IX, a divalent, cation-independent antibody, recognizes factor IX when used on immunoblots of non-denatured, non-reduced human plasma.

Factor IX is a 55 kDa, single chain, vitamin K-dependent plasma zymogen which plays a key role in the intrinsic and extrinsic blood coagulation systems. Hereditary deficiencies or dysfunctions of factor IX cause hemophilia B or “Christmas Disease” (the surname of the first family described).

A disulfide bond in factor IX connects the N-terminal sequence (light chain) of factor IX to the C-terminal sequence (heavy chain). Upon activation of factor IX to factor IXa by factor Xla in the intrinsic system, an 11 kDa activation peptide is removed from the factor IX molecule by cleavage of two peptide bonds. These changes allow the exposure of the serine protease site on the heavy chain which can then activate factor X in the presence of factor VIII, Ca\(^{2+}\), and phospholipid.

Factor IX can be similarly activated by the extrinsic system, (i.e., the tissue factor-factor VII complex,)\(^1\) Factor IX is synthesized in liver parenchymal cells and requires a post-translational, vitamin K-dependent, modification in order to become a mature plasma zymogen. When patients lack Vitamin K, or take oral anticoagulants that interfere with the metabolism of vitamin K, a hypocoagulable or antithrombotic state is induced. This state stems from the diminished ability of factor IX to bind to phospholipids.

Factor IX concentration in human plasma ranges between 2.5-5 µg/ml and its half life is ~24 hours. Human factor IX gene is about 40 kb in size and is localized at the distal end of the X-chromosome. The gene has been completely sequenced\(^2\) and so far more than 50 gross or subtle mutations have been discovered.\(^3\)

Assays of factor IX antigen levels are useful for:
1. Initial characterization of the genetic defect in patients affected by hemophilia B.
2. Detection of female carriers of hemophilia B in families affected by mutant genes that are expressed by dysfunctional factor IX.
3. Prenatal diagnosis by fetal blood sampling when molecular genetic techniques cannot be used.
4. In vitro studies of the role of factor IX in the intrinsic or extrinsic pathways of blood coagulation.

The antibody is useful for the preparation of factor IX depleted plasma and for purification of factor IX.

Reagent
The product is provided as purified antibody in 10 mM HEPES, 140 mM NaCl, pH 7.4, containing 0.05% sodium azide as a preservative.

Precautions and Disclaimer
This product is for R&D use only, not for drug, household, or other uses. Please consult the Safety Data Sheet for information regarding hazards and safe handling practices.

Storage/Stability
For continuous use, store at 2–8 °C for up to one month. For extended storage, freeze in working aliquots. Repeated freezing and thawing is not recommended. Storage in "frost-free" freezers is not recommended. If slight turbidity occurs upon prolonged storage, clarify the solution by centrifugation before use.
Product Profile
At a concentration of 5.0 μg/ml, the antibody inhibits >90% factor IX activity in human plasma as measured by the Activated Partial Thromboplastin Time (APTT) assay.

References