SUMMARY

One of the limitations of many bridging experiments in neural transplantation is that the CNS tissues cannot be sutured. Fibrin glue is a two-component system derived from whole blood which, when mixed, reproduces the final stage of blood coagulation and solidifies. Many experimental studies of humans and animals show that fibrin glue repair of peripheral nerves is almost equivalent to microsurgical sutures. In this study, we attempted to extend its use to CNS tissues and transplants. Two techniques were tried: (1) Bilateral parietal knife cuts were performed by stereotaxic technique in six rats. Fibrin glue was applied in the right-side cortical lesion. Immunohistochemistry using antisera to tyrosine hydroxylase (TH), glial fibrillary acidic protein (GFAP), laminin and neurofilament (NF) was essentially similar between the control and treatment groups. The immunoreactivity of each marker revealed no significant differences between the two groups on days 1, 7 and 30. There was no difference in terms of gliosis or microvascular proliferation. (2) Embryonic day 16 fetal locus coeruleus was grafted together with E16 cortex to the anterior chamber of sympathectomised eyes. In the six eyes of the glue treatment group, the parietal cortical piece and the locus coeruleus piece were joined together before grafting by immersing them in the solution of fibrin glue. In the eight eyes of the control group, pieces of parietal cortex and locus coeruleus were introduced individually and approximated by gently pressing the cornea. The sizes of double grafts showed no significant difference between groups during six weeks postgrafting. The immunohistochemical pictures using antisera against TH, GFAP and laminin were similar in both groups. Catecholaminergic fibers from the grafted locus coeruleus were found bridging over into the parietal cortical piece in both the control and treatment groups. There was no significant difference in TH-positive nerve fiber density between tissue glue-joined and control double intraocular grafts. In conclusion, fibrin glue can be used as an adhesive agent in CNS tissues without hampering the outgrowth of neurites or causing adverse tissue reactions in fetal or adult nervous tissues.

KEY WORDS

fibrin glue, CNS repair, brain development, transplantation, locus coeruleus, cortex cerebri

INTRODUCTION

In the intensive search for methods to improve the regenerative capability of CNS tissues, many different bridging models have been attempted. One of the limitations of such experiments is that CNS tissues cannot be sutured. The CNS microvasculature cannot sustain suturing and the fragility of the tissue makes attempts to use stitches hazardous or useless. One approach uses transplants which may serve as bridges for regenerating axons across a site of injury /25/. Alternative methods for bridging by inserting peripheral nerve pieces, or lodging tissues into the CNS, or anchoring to the dura matter with microsutures could provide moderate but not strong joining.

For peripheral nerve repair, several sutureless methods have been developed, including fibrin glue
Fibrin glue, or tissue glue, is a two-component system derived from whole blood which, when mixed, reproduces the final stage of blood coagulation to form a viscous adhesive. This glue is commercially available and used both clinically and experimentally in Europe and Japan. It consists of highly concentrated human fibrinogen which, after adding aprotinin, reacts with the calcium-activated thrombin to form a sealant that can maintain tissue approximation. Most of the experimental studies on humans or animals have shown that fibrin glue repair of peripheral nerves was equivalent to microsurgical sutures. Its major advantage over sutures is that the time required for the procedure is shorter and the anastomosis is easier to perform, especially in hard-to-access areas.

In this study, we attempted to extend the use of fibrin glue to CNS tissues and transplants. First, a rodent CNS lesion model was used to evaluate any adverse effect of the glue in lesioneon CNS areas. Next, we used the fibrin glue as an adhesive agent between two pieces of intraocular fetal grafts (locus coeruleus and cerebral cortex) to evaluate its influence on survival and growth of fetal tissue grafts. Finally, the intraocular double graft model allowed us to determine whether nerve fibers could bridge a layer of fibrin glue at the interface between two brain tissue areas.

MATERIALS AND METHODS

Fibrin glue

The fibrin glue used for the present study was a fibrinogen-based compound with double sealant components (Beriplast® P, Behring, Behringwerke AG, Marburg, Germany). The 1 ml Beriplast® P set contains:

Vial 1: Fibrinogen concentrate consisting of 115-232 mg dry substance, containing a human plasma protein fraction with 65-115 mg fibrinogen and a human plasma protein fraction with a factor XIII activity of 40-80 U.

Vial 2: Aprotinin solution consisting of 1 ml solution containing 1000 KIU of bovine lung aprotinin.

Vial 3: Thrombin consisting of 4.9-11.1 mg dry substance containing a human plasma protein fraction with a thrombin activity of 400-600 IU.

Vial 4: Calcium chloride solution consisting of 2.5 ml solution containing 14.7 mg calcium chloride-2H₂O (40 mmol/l).

Before application, the two sealant elements were prepared separately by transferring the aprotinin solution to the fibrinogen concentrate and transferring the calcium chloride solution to the thrombin. To apply the fibrin glue, a tuberculin syringe was filled with each sealant element. By pushing the two syringe plungers simultaneously, a coagulating glue forms between the tips.

Fibrin glue in a CNS lesion model

Adult 150-g female rats (Sprague-Dawley, B&K Universal, Stockholm) were used to study the effect of fibrin glue in the CNS lesion model. Under general halothane anesthesia, the animal was mounted in a stereotaxic frame. A sharp blade with a flat cutting edge, 2 mm in length, was clamped onto the frame. Bilateral parietal craniectomies were performed to expose 3 x 3 mm² areas on both sides. The central bony ridge over the sagittal sinus was preserved to impede the liquid flow from one side to the other. Under stereotaxic guidance, the centers of the craniectomies were identified for cortical incision (AP 1.5, L/R 1.5, V 0 from bregma). The dura was opened and bilateral cortical incisions were performed by moving the clamped blade 3 mm downward (from V 0 to V 3, see Fig. 1). The confronting vessels on the cortical surface were cauterized and severed before cutting to prevent profuse bleeding. Fibrin glue was applied into the right-side cortical lesion. Care was taken to avoid any fluid leak to the left side. After the glue hardened, the craniectomies were covered by pieces of spongostan and the wound closed.

The six rats were grouped into three groups and sacrificed on days 1, 7 and 28 respectively. The animals were anesthetized with sodium pentobarbital (Mebumal 40 mg/kg i.p.) and perfused via the ascending aorta with 50 ml calcium-free Tyrode's solution followed by 50 ml formalin-picric acid mixture (4% paraformaldehyde, 0.4% picric acid in 0.16 M phosphate buffer, pH 7.4). Brains were removed and processed for indirect immunohistochemistry using antibodies against tyrosine hydro-
xylase (TH), glial fibrillary acidic protein (GFAP), laminin, and neurofilament protein (NF). Tissues were examined and photographed using epifluorescence microscopy (Nikon-Microphot). Densities of TH, GFAP, laminin and NF immunoreactivities were semiquantitatively scored on all sections. Averages from 6-10 sections from a given brain were then calculated. A semiquantitative scale from 0-5 was used.

Fibrin glue in double intraocular grafts (locus coeruleus + cortex cerebri)

Young adult 150-g female rats (Sprague-Dawley, B&K Universal, Stockholm) were used as hosts to study the effect of fibrin glue in double intraocular grafts. All eyes were sympathetically denervated by superior cervical ganglionectomy one week before transplantation. Fetuses from pregnant
Fig. 2: Immunohistochemistry of lesioned cerebral cortices on days 1, 7 and 30. The histologic pictures with four different immunoreactive markers were similar between the control (right side) and the glue-treated (left side) cortices. GFAP: glial fibrillary acidic protein, LAM: laminin, NF: neurofilament protein, TH: tyrosine hydroxylase. a: day 1; b, d and f: day 7; c, e and g: day 30. Scale bar = 500 μm in LAM. Scale bar = 100 μm in GFAP and NF.
rats of the same strain were used as donors of CNS tissue grafts. Pieces of parietal cortex and locus coeruleus from E16 fetal brains were dissected out. In the treatment group, the cortical and locus coeruleus pieces were joined together by immersing them in the fibrin glue solution. The superfluous glue around the joined piece was trimmed after the glue had hardened. After pretreatment of all eyes with a drop of 1% atropine solution, intraocular grafting was performed under ether anesthesia. Grafts were introduced through a tiny opening of the cornea as previously described/19-21/ using a modified Pasteur pipette. The fibrin glue-treated combined graft was introduced as a single piece, while in the control group pieces of parietal cortex and locus coeruleus were introduced individually and juxtaposed by gentle pressure on the cornea. After grafting, the volumes of the transplants were measured at regular intervals by stereomicroscopy. The graft size was estimated by taking the longest diameter of the graft multiplied by the diameter perpendicular to it /4/. Eleven animals were used which consisted of six eyes in the treated group, eight eyes in the untreated group and six eyes of cortico-cortical glue-treated double grafts as negative controls (see Fig. 1).

The animals were perfused six weeks after grafting using procedures as stated above and processed with the same methods for indirect immunohistochemistry using antibodies against TH, GFAP and laminin. Scoring of TH-immunoreactive nerve fiber density, GFAP-like, and laminin immunoreactivities was performed on all sections. Averages from 6-10 sections from a given graft were then calculated to generate means of TH-immunoreactive nerve fiber density, GFAP-like and laminin immunoreactivity per section. A semiquantitative scale from 0-5 was used.

Statistical analysis

Fibrin glue in CNS lesion model

The data were analyzed by nonparametric Mann-Whitney tests. Within each group, the data from the immunoreactivity of glue-applied sides were compared with the data from the corresponding immunoreactivity of control sides.

Fibrin glue in double intraocular grafts (locus coeruleus + cortex cerebri)

Graft sizes were analyzed by single-factor repeated measures ANOVA test. The univariate analysis was based on repeated measurements on the same rats. Number of days after surgery and number of eyes were treated as within-subject factors. Rats were grouped within the treatment groups. The type of treatment was a between-subject factor. The measurements of TH-immunoreactive nerve fiber density, GFAP-like and laminin immunoreactivity were analyzed by ANOVA.

RESULTS

Fibrin glue in CNS lesion model

The right parietal cortex cuts which received fibrin glue were compared with the left parietal cortices that received lesions only. The patterns of immunohistochemistry using antisera against TH, GFAP, laminin and NF were similar between the control and treatment groups on days 1, 7 and 30 (Fig. 2). There was no difference in the degree of gliosis or microvascular proliferation. No morphologic evidence of any special tissue reaction could be found in the treatment group. Using the semiquantitative data, none of the immunoreactive markers revealed in the glue group differed significantly from the controls on days 1, 7 or 30 (Fig. 3).

Fibrin glue in double intraocular grafts (locus coeruleus + cortex cerebri)

Growth of double intraocular grafts

With or without fibrin glue, joined fetal (E16) locus coeruleus and cortex cerebri double pieces were grafted to the anterior chamber of the eye of sympathetically denervated host rats. All but two of the grafts (in the group of cortico-cortical grafts) survived well and became vascularized from the host iris. As shown in Fig. 4, there was no significant difference in volume change between the glue-treated and untreated locus coeruleus plus cortex cerebri grafts (P = 0.25 on day 5, P = 0.38 on day 14, and P = 0.52 on day 25, ANOVA).
Fig. 3: Comparison of the immunoreactivity between glue-treated and control lesioned cortex with four immunohistochemical markers based on 6-10 serial sections from each brain (n = 2 for each group). The density of neural, glial and vascular elements with four immunohistochemical stains was similar in the glue-treated and control lesioned cortex on days 1, 7 and 30. The fiber density of each section was graded from 0 to 5. LAM: laminin, GFAP: glial fibrillary acidic protein, NF: neurofilament protein, TH: tyrosine hydroxylase.

Fig. 4: Volume of double intraocular grafts (locus coeruleus + cortex cerebri, E16) on days 5, 14 and 25. No significant difference in volume change was noted between the glue-treated and untreated locus coeruleus plus cortex cerebri grafts (glue-treated double grafts, n=6; untreated controls, n=8; ANOVA).

Effects of fibrin glue on TH-positive nerve fiber density, GFAP and laminin immunoreactivity in intraocular double grafts

The catecholaminergic fibers in the intraocular double grafts were examined using TH immunohistochemistry. There was no significant difference in TH-positive nerve fiber density between glue-joined and untreated locus coeruleus plus cortex cerebri grafts (Figs. 5, 6; P = 0.76, ANOVA). Comparing the TH-positive fiber density in the above glue-treated or untreated locus coeruleus plus cortex cerebri grafts with the glue-treated cortico-cortical grafts, significant differences were noted (P = 0.017 for the untreated and cortico-cortical grafts, P = 0.01 for the glue treated and cortico-cortical grafts, ANOVA; Fig. 5). Studies of GFAP and laminin immunoreactivity in the grafts revealed no obvious differences between the different treatments (see also Fig. 5).

DISCUSSION

The fact that no morphologic difference could be found between the fibrin glue-treated and control lesioned cortex suggests that fibrin glue does not produce harmful effects in CNS tissue. Gliosis and
microvascular proliferation did not increase in the glue-treated lesioned cortices in the early, intermediate or late periods after injury. Studies of TH and NF immunoreactivity also indicated that neural components of the CNS tissue are unaltered after the application of glue. As shown by many authors /1-3,8-11,13,14,16,22,24,26,28/, fibrin glue produces no adverse tissue reaction in PNS tissues. After axotomy, axons can penetrate the glue region and grow in the distal stumps.

The major controversy concerning the application of fibrin glue in peripheral nerve repair is whether it can replace the conventional microsuture technique. Studies by several authors /3,9,10,14,22/ have shown that the effect of fibrin glue in nerve coaptation is comparable to microsutures in tensile strength, regenerative axonal counts and, although slower in conduction velocity, electrophysiologic responses. The advantages of fibrin glue in surgery are rapidity of application, good hemostasis, easier stabilization of small grafts and possibility of anastomosis in some difficult anatomical positions /8,15,22/. The disadvantage of possible bacteria and virus transmission has been resolved by inactivation. Screening for hepatitis B antibodies since the early 1980s and for AIDS virus antibodies since the 1990s has been done to scrutinize both source material and the final product /15/. With a relatively aseptic technique, as presented in this study, infection is not a problem with the use of fibrin glue in the CNS tissues.

The vulnerable nature of the microvasculature of CNS tissue, the fragile texture of the parenchyma and the sophisticated neural networks combine to make suturing impossible for reparative intervention. The present intraocular double grafting experiments have demonstrated a possible role for fibrin glue to act as a CNS adhesive agent to glue CNS tissues together. The glue-joined fetal double grafts survived and grew on the iris. Statistical

Fig. 5: Immunohistochemistry of double intraocular grafts 37 days postgrafting. The TH-positive nerve fiber density was similar between glue-joined (upper) and untreated (lower) locus coeruleus plus cortex cerebri grafts. Scale bar = 200 μm.
Comparison of TH-positive nerve fiber density, GFAP-like and lamin immunoreactivity between glue-joined double intraocular grafts (G: locus coeruleus + cortex cerebri, E16, n=6) and untreated double intraocular grafts (C: locus coeruleus + cortex cerebri in HBSS, E16, n=8). Double pieces of cortex cerebri (Co + Co: cortex cerebri + cortex cerebri, E16, n=4), joined by fibrin glue, were also used as negative controls. There is no significant difference in TH-positive nerve fiber density between tissue glue-joined and untreated double intraocular grafts. Significant differences are noted between the above two groups and the cortico-cortical grafts (P=0.0165 for C; Co + Co, P=0.0122 for G, Co + Co, ANOVA). No difference is seen in the density of GFAP-like and lamin immunoreactivity within the three groups (ANOVA).

Fig. 6: Comparison of TH-positive nerve fiber density, GFAP-like and lamin immunoreactivity between glue-joined double intraocular grafts (G: locus coeruleus + cortex cerebri, E16, n=6) and untreated double intraocular grafts (C: locus coeruleus + cortex cerebri in HBSS, E16, n=8). Double pieces of cortex cerebri (Co + Co: cortex cerebri + cortex cerebri, E16, n=4), joined by fibrin glue, were also used as negative controls. There is no significant difference in TH-positive nerve fiber density between tissue glue-joined and untreated double intraocular grafts. Significant differences are noted between the above two groups and the cortico-cortical grafts (P=0.0165 for C; Co + Co, P=0.0122 for G, Co + Co, ANOVA). No difference is seen in the density of GFAP-like and lamin immunoreactivity within the three groups (ANOVA).

analysis showed no significant difference between the two groups regarding graft volumes. The study of TH-positive nerve fiber densities of the intraocular double grafts six weeks after grafting revealed no significant difference between the glue-treated and the untreated groups. Likewise, GFAP and lamin immunoreactivity also appeared similar. Since TH-positive fibers found in the locus coeruleus plus cortex cerebri grafts could not be derived from the sympathetomized iris, their presence in the cortex part of the double graft proves that a coeruleo-cortical TH-positive pathway had formed. Clearly, the fibrin glue did not interfere with the growth of nerve fibers; nor did it cause any untoward gliosis or disturbed microcirculation in the grafts.

Study of the ultrastructure of fibrin glue with confocal laser 3D microscopy, liquid permeation and turbidity /5/ shows that the glue from normal fibrinogen is composed of straight rod-like fiber elements which sometimes originate from denser nodes. The tautness and porosity of the glue networks can be altered by changing the concentration of thrombin and fibrinogen. Thus increasing either thrombin or fibrinogen concentrations will cause the glue networks to become tighter and the porosity to be decreased. The fiber strands also become shorter. Glue porosity of the network can also be modified by changing the ionic strengths, coexisting proteins (e.g. albumin, lipoprotein), dextran, etc.

For the catecholaminergic cells in the locus coeruleus graft to survive and grow, nutrients and diffusible factors (such as trophic factors, from either the iris or the co-grafted cortical piece) must be able to reach the cells in the graft /18-21/. The endothelial cells from the iris must also have access to the graft, otherwise the graft would degenerate and die. The present study supports the finding that nutrients and trophic factors can pass through the fibrin glue networks to reach the graft. It means, with the present formula, the formed fibrin polymers create a mesh that will hold two graft pieces together but also permit the transmission of essential molecules through its pores. The endothelial cells from the iris can penetrate the fibrin meshwork around the graft and finally establish the microvasculature. Likewise, TH-positive fibers from the locus coeruleus can also penetrate the fibrin glue in between the two grafts and reach the cortical piece. The fibrin meshwork is known to be absorbed by the fibrinolytic system (e.g. plasmin) one week after application. The feasibility for the lysis of fibrin glue can also be altered by structural modification of the extended dimeric multidomain fibrin molecule (e.g. as shown in the studies of Blombäck et al. and Nossel & Kaplan /5,17/, a scheme involving fibrin I and fibrin II, in which the former is the result of releasing the fibrinopeptide only. Fibrin II lacks both fibrinopeptides A and B and is thought to be involved in lateral anchoring; as a consequence it is more resistant to fibrinolysis), presence of physiological stabilizing factor XIIIa, interactions between plasminogen, thrombin and fibrin, cross-linked fibronectin in the tissues, platelets, etc. /5,7,17,25/.
In the anterior chamber of the eye, the fibrin glue is exposed to the chamber fluid, the iris, locus coeruleus and cortical graft. Since there was no significant difference in graft growth or final sizes between glue-treated and control double grafts, it appears as if the glue coat did not significantly impair the rate of formation or amount of vascular connections between host iris and grafts. The lysis of the glue will be affected by numerous factors and interactions. The timing of the penetration of endothelial cells and neurites, in relation to fibrinolysis, remains to be further analyzed.

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