Microcontact Printing of Cholinergic Neurons in Organotypic Brain Slices

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Alzheimer's disease is a severe neurodegenerative disorder of the brain, characterized by beta-amyloid plaques, tau pathology, and cell death of cholinergic neurons, resulting in loss of memory. The reasons for the damage of the cholinergic neurons are not clear, but the nerve growth factor (NGF) is the most potent trophic factor to support the survival of these neurons. In the present study we aim to microprint NGF onto semipermeable 0.4 µm pore membranes and couple them with organotypic brain slices of the basal nucleus of Meynert and to characterize neuronal survival and axonal growth. The brain slices were prepared from postnatal day 10 wildtype mice (C57BL6), cultured on membranes for 2–6 weeks, stained, and characterized for choline acetyltransferase (ChAT). The NGF was microcontact printed in 28 lines, each with 35 µm width, 35 µm space between them, and with a length of 8 mm. As NGF alone could not be printed on the membranes, NGF was embedded into collagen hydrogels and the brain slices were placed at the center of the microprints and the cholinergic neurons that survived. The ChAT+ processes were found to grow along with the NGF microcontact prints, but cells also migrated. Within the brain slices, some form of re-organization along the NGF microcontact prints occurred, especially the glial fibrillary acidic protein (GFAP)+ astrocytes. In conclusion, we provided a novel innovative microcontact printing technique on semipermeable membranes which can be coupled with brain slices. Collagen was used as a loading substance and allowed the microcontact printing of nearly any protein of interest.

Keywords: cholinergic neurons, microcontact printing, nerve growth factor, organotypic brain slices, brain-on-a-chip

INTRODUCTION

Alzheimer's disease is a neurodegenerative disorder leading to a progressive decline in cognitive and intellectual functions without a clear causative event. The cardinal pathological features of Alzheimer's disease (AD) are extracellular β-amyloid (Aβ) depositions (plaques) and intraneuronal hyperphosphorylated tau inclusions (neurofibrillary tangles (NFTs)). This is accompanied by chronic inflammation as a contributor to the neurodegenerative processes observed in AD but also in PD (1, 2). There is only a weak correlation between cognitive decline, Aβ plaques, and NFTs, but the density of neocortical synapses strongly correlates with all three, indicating that synaptic loss is the major correlate of cognitive impairment in AD (3). The main populations of cholinergic neurons are located in the basal forebrain: the nucleus basalis of Meynert (nBM) and medial septum. They provide projections to the entire neocortex and hippocampus, synthesizing and releasing...
acetylcholine (4, 5). It has been demonstrated that the selective
degeneration of this transmitter-specific neuronal population in
the nBM is a major hallmark in AD (6, 7). Indeed, the decline
of cholinergic neurons directly correlates with a decrease in
cognitive and intellectual functions (8). However, so far, it is still
unclear if the cell death of cholinergic neurons is a primary event
in AD or is caused by the dramatic deposition of Aβ plaques or
tau NFTs in the cortex and hippocampus.

The survival of basal forebrain cholinergic neurons is
dependent on the classical nerve growth factor (NGF) (9–
11). The NGF is synthesized in the target area of cholinergic
neurons: the cortex and hippocampus. As NGF is a target-
derived neurotrophic factor, it is endocytosed by the cholinergic
nerve fibers and retrogradely transported to the somata in the
nBM/septum where an NGF-dependent transcriptional program
is activated (12). Two different cell-surface receptors, as well
as the state of NGF (unprocessed pro-form or mature form),
determine the activity and function of NGF. Mature NGF has
a higher affinity for the tropomyosin receptor kinase A (trkA)
receptor promoting survival and growth, whereas pro-NGF preferentially binds to neurotrophin receptor p75NTR-mediated
apoptotic signaling (13). Increasing evidence supports that the
NGF function is affected in AD (14) as an imbalance between the
NGF/TrkA-mediated survival signaling and pro-NGF/p75NTR-
mediated apoptotic signaling may occur. Moreover, the TrkA-
dependent retrograde transport of NGF could be impaired for
the Aβ and tau pathology and cholinergic neurons are not able
to take full advantage of the NGF (12). Thus, NGF may also have
potential therapeutic implications in AD, as therapeutic strategies
aim to deliver NGF directly into the brain (15–17).

Organotypic brain slice cultures bridge the gap between in
vitro cell cultures and in vivo animal experiments. In contrast
to homogenous single cell cultures, in organotypic brain slice
cultures, the complex three-dimensional architecture of the brain
is preserved, simulating more in vivo-like situations (18–20).
Additionally, organotypic brain slice cultures permit markedly
reducing the number of animal experiments. As the cell death
of cholinergic neurons is the central hallmark of AD, organotypic
brain slices are a potent tool to study the neurodegeneration of
cholinergic neurons in AD. The presence of cholinergic neurons
in organotypic slices is verified in the septum/hippocampus (4)
and the nBM, the latter extensively studied in our lab (9, 21–
24). It is well-established that NGF is required to maintain
cholinergic neurons in organotypic brain slices (9, 11, 21,
25). In this study, NGF supports the survival of cholinergic
neurons in nBM organotypic brain slices when applied to a
medium (9, 21). We also developed a model, where NGF was
locally applied directly onto brain slices using collagen hydrogels
(1, 11, 26).

To take it a step further, we aimed to immobilize the
growth factors in a pattern next to the slice enabling guided
nerve fiber outgrowth. Therefore, we took advantage of a
high-resolution (sub-μm range) patterning technology called
microcontact printing (μCP). The μCP technique is referred to
as soft lithography and is widely used to immobilize proteins onto a variety of different background materials (27,
28). However, μCP seems to be a straightforward technique
for printing clear and even plasma-activated surfaces, but
this does not apply to pored membranes. Currently, only a
single study describes printing onto membranes using heavy-
ion etched polycarbonate membranes (29). Printing onto
pored membrane inserts, however, has not been demonstrated
so far. Recently, we were successful in printing antibodies
onto pored membranes and coupled these antibodies with the
growth factor gial-cell line-derived neurotrophic factor
(GDNF) to stimulate the nerve fiber growth of dopaminergic
neurons (1).

The present study aimed to establish the microcontact
printing of the protein NGF on semipermeable 0.4 μm
pore membranes and to couple them with organotypic
tissue slices of the nBM. To print the NGF, collagen was
used as a loading biomaterial. We will demonstrate that
cholinergic neurons will survive and grow along the NGF
microcontact prints.

MATERIALS AND METHODS
Organotypic Brain Slices of the nBM
Organotypic chopper brain slices were prepared as reported in
detail in our lab (21). Briefly, postnatal day 8–10 C57BL/6 wildtype mouse pups were rapidly decapitated and their brains
were dissected under sterile conditions. The nBM was dissected
according to our scheme (21) and 300 μm slices were chopped
on a McIlwain Tissue Chopper (Mickle Laboratory Engineeringe
Co. LTD, Loughborough, England). The brain slices were
carefully transferred to an Isopore™ 0.4 μm pore PC membrane
(HTTP02500, Merck Millipore, Burlington, Massachusetts,
United States) with μCP lanes (see below). These membranes
were transferred onto semipermeable 0.4 μm pore cell culture
inserts (PICM03050, Merck Millipore) which were placed into
6-well plates (Greiner Bio-One, Kremsmünster, Austria). Each
well-contained 1.1 ml of sterile-filtered culture medium (50%
MEM/HEPES (Gibco, Carlsbad, California, United States), 25%
heat-inactivated horse serum (Gibco/Lifetech, Austria), 25%
Hanks’ solution (Gibco), 2 mM NaHCO3 (Merck, Austria), 6.5
mg/ml glucose (Merck, Germany), and 2 mM glutamine (Merck,
Germany), at pH 7.2)). The brain slices were cultured at 37°C
with 5% carbon dioxide (CO2) for 2–6 weeks and the culture
medium was changed once a week. The slices attached to the
membranes were flattened and became transparent. The brain
slices were cultured with or without 100 ng/ml of recombinant
anti-mouse β-NGF (50385-MNAC, Sino Biological, Germany)
in a culture medium. After 2–6 weeks, the slices were fixed
for 3 h at 4°C in 4% paraformaldehyde and stored at 4°C in
10 mM of phosphate-buffered saline (PBS) until use. All the
experiments conformed to the Austrian guidelines on the ethical
use of animals and were in line with the reduce, refine and
replace (3Rs) rule as all efforts were made to reduce the number
of animals. In fact, from one mouse pup, we were able to
generate 50–100 brain slices depending on the area and purpose.
All animal experiments were defined as “Organenentnahme”
according to the Austrian laws. For this project, 47 mouse pups
were used.
Preparation of Collagen Hydrogel Solution

Collagen hydrogels were prepared as we have described in detail (11). As a crosslinker, 4S-Star-polyethylene glycol succinimidyl succinate (4S-StarPEG) (JKA7006-1G, Sigma, St. Louis, Missouri, United States) was used. Two mg/ml sterile bovine collagen solution type I (804592-20ML, Sigma) was linked with 0.4 mM of 4S-StarPEG in PBS at pH 7.4. The Collagen-polyethylene glycol hydrogel (PEG) solution was loaded with a recombinant anti-mouse β-NGF (50385-MNAC; Sino Biological) in a final concentration of 10 ng/µl of NGF or fluorescent Alexa-546 anti-goat antibodies (final concentration: 20 ng/µl). An equal volume of PBS was added to generate the control collagen hydrogel microcontact prints. During the handling, all the components were kept on ice to prevent pre-mature gel formation. Approximately 100 µl of collagen hydrogel ink solution (≈ 1 µg NGF) was immediately applied onto the µCP stamp (see below).

Microcontact Printing

The silicon wafer master mold (Figure 1A) was a kind gift from Jenny Emmnäus and Janko Kajtaz (Department of Biotechnology and Biomedicine, DTU Bioengineering, Technical University of Denmark), and has been used previously in a common publication (1). The master mold has been fabricated by photolithography using a silicon oxide layer (4.7 µm) and is described elsewhere.

The Micropatterned stamp fabrication was performed as we have previously described (1). Polydimethylsiloxane
is highly suitable for protein adsorption and transfer due to its hydrophobic surface. The polydimethylsiloxane (PDMS) prepolymer (SylgardTM 184 Silicone elastomer kit, 001004176976, Dow) arrived in two components. The elastomer curing agent was carefully mixed with an elastomer base solution in a concentration of 1:10. The surface relief of the PDMS stamps was formed by casting and curing liquid PDMS against the micropatterned silicon wafer master mold (Figures 1B,C). The raised and lowered regions of the silicon wafer were mirrored into the stamps and the final patterns were defined. After being left to cure overnight at 60°C, the solid PDMS was peeled off the mold and the stamps were cut to size with a scalpel for further use.

The μCP was performed similarly as reported from our lab (1) but modified. Microcontact printing uses an elastomer stamp which adsorbs the “ink” solution and transfers it to a surface at a very high resolution. Approximately 100 μL of the liquid collagen hydrogel ink solution (loaded with PBS, NGF, or anti-goat Alexa-546 antibodies) was applied directly onto the micropatterned stamp. To distribute the ink solution equally, a coverslip was placed on top. After 15 min of incubation at 37°C, the coverslip was removed and used to carefully strike off the remaining ink solution, once with and once against the lanes of the pattern. The excess solution on the borders of the pattern was removed using filter paper without touching the printing surface and was left to air dry for a minute. As soon as the stamp was completely dry, the ink solution was transferred to the semipermeable membrane by pressing it on with an 18 grams weight for 60 min at room temperature. The position of the stamp was marked with four small dots of permanent marker for the arrangement of the slices. Then, the weight and the stamp were carefully removed from one corner. The membranes were sterilized under a UV light for 20 min, equilibrated with the slice medium, and placed on the inserts before arranging the brain slices. To optimize the insert before arranging the brain slices. To optimize the μCP experiments, we used the Isopore™ 0.4 μm pore PC membrane (HTTP02500, Merck Millipore). We also compared μCP on Omnipore™ 0.45 μm PTFE membranes (JHPW02500, Merck Millipore) and LCR 0.45 μm PTFE membranes (FHL02500, Merck Millipore).

**Immunohistochemistry**

Immunohistochemistry was performed as previously described under free-floating conditions (21, 30). This method allows the antibody to penetrate from both sides during incubation, enhancing the sensitivity of the staining. The outgrowth processes were considered to be the single-cell layers on the membrane. First, the fixed brain slices were incubated in 0.1% Triton-PBS (T-PBS) for 30 min at room temperature with soft shaking. After incubation, the brain slices were washed 3 × 3 min with PBS and subsequently blocked in 20% horse serum/0.2% bovine serum albumin (BSA)/T-PBS for 30 min at room temperature with soft shaking. Following the blocking, the brain slices were incubated in 0.2% BSA/T-PBS with primary antibodies, namely, choline acetyltransferase (ChAT) (Merck AB144P, 1:750), glial fibrillary acidic protein (GFAP) (Merck AB5541, 1:2000), laminin (Sigma L9393; 1:500), p75NTR NGF receptor (Abcam ab52987; 1:750), microglial Iba-1 (Wako 019-19741; 1:500), microtubule-associated protein-2 (MAP-2) (Chemicon MAB3418; 1:500), or NGF (Cedarlane MC51, 1:250) for 48 h at 4°C. After incubation, the brain slices were washed 3 × 3 min with PBS and incubated with the corresponding green fluorescent Alexa-488 (or red fluorescent Alexa-546) secondary antibodies (1:400 in 0.2% BSA/T-PBS) for 1 h at room temperature while shaking. The secondary antibodies were: anti-goat for ChAT, anti-chicken for GFAP, anti-rabbit for laminin, p75NTR, NGF, Iba-1, and anti-mouse for MAP-2. The brain slices were counterstained with the blue fluorescent nuclear dye, 4,6-diamidino-2-phenylindole (DAPI) (1:10,000 diluted in T-PBS), for 30 min. The brain slices were washed again with PBS before being mounted on glass slides with Mowiol (Carl Roth, Karlsruhe, Germany). The staining was visualized with a fluorescence microscope (Olympus BX61, Olympus Corporation, Shinjuku City, Tokyo, Japan) and Openlab software 5.5.0 (Improvision, Germany). For the co-stainings, the slices were washed 3 × 5 min with PBS before the application of the second primary antibody. Some sections were stained with the chromogenic DAB.

**Data Analysis and Statistics**

Quantitative analysis was performed blinded under the microscope. The ChAT+ nBM neurons were recognized when clear cytoplasmic staining with at least one neuronal fiber extension and a definable nucleus was observed. The slices were excluded if the number of ChAT+ neurons was <20/slice. As two chopper slices were placed side by side in the center of the μCP, the outgrowths projecting upwards and downwards were separately quantified. The number of ChAT+ processes was evaluated along the NGF μCP and they were counted as single nerve fibers, nerve fiber bundles, thick nerve fiber networks, and outgrowths with and without ChAT+ cells. To evaluate the length of the outgrowth, a picture was taken under the Olympus BX61 fluorescence microscope and the pixel number of the outgrowth was measured using the OpenLab 4.0.4 software connected to a MAC computer (Apple, Cupertino, California, United States). The pixel number was measured from the slice border to the very top of the outgrowth. The values obtained were then converted to micrometers based on a scale bar. To evaluate the intensity of the NGF μCP, the optical density (OD) was measured. Therefore, a picture was taken with 1 s exposure time and 10×-magnification and afterward was converted into greyscale. Then a rectangle of 30 × 100 pixels was chosen within a printed lane and transferred to Photoshop Elements 2.0 (Adobe, San Jose, California, United States). After inverting the picture, the OD mean was taken from the histogram and corrected for the background. The OD measurement was repeated 3 times per print. The sample size (n) pertains to the number of analyzed mice. All the values are given as mean ± SEM. Statistical analysis was performed by one-way ANOVA with a subsequent Fisher least significant difference (LSD) post-hoc test, where p < 0.05 represents significance.
**FIGURE 2** | Characterization of nerve growth factor (NGF) µGP on semipermeable membranes. Printing of collagen alone (A) or NGF alone (B) does not show any positive staining, however, NGF loaded into collagen hydrogel solution shows a typical printing when stained with Alexa-546 anti-NGF antibodies (C). The staining is specific, as it is not seen in the green channel (D). A concentration experiment shows that printing with 1,000 ng NGF per stamp (E) gives the best results, while 100 ng (F) and 10 ng (G) NGF do not show good printing. The stability of the prints (1,000 ng NGF loaded into collagen hydrogel solution) in the medium is shown in (H–J), with strong bands after printing [day 0, (H)], no change after 9 days in the medium (I), and a significant decrease after 14 days in the medium (J). The values in (E–J) give the optical density of the printed pattern from 0 (white) to 255 (black) corrected for the background and measured by computer-assisted imaging. The values are presented as mean ± SEM, with the number of independent experiments (n) and statistical differences (*p < 0.05; ***p < 0.001) with ANOVA and Fisher least significant difference (LSD) post hoc test. Scale bar in A = 175 µm (A–D), 102 µm (E–G), 175 µm (H–J).
RESULTS

Characterization of $\mu$CP Using Antibodies

To characterize the method, we first performed $\mu$CP on a fluorescent antibody, as established in our lab. Using the wafer master mold (Figure 1A), PDMS stamps (Figure 1B) were produced with a size of 2 x 8 mm (Figures 1C,D) and 28 lanes (Figure 1D). The printing of an anti-goat Alexa-546 antibody showed several red lines under the fluorescence microscope (Figure 1E), with a line width of 30 µm and a space of 30 µm (Figure 1F). The staining was not seen in the green channel, showing the specificity of $\mu$CP (Figure 1G).

Characterization of NGF $\mu$CP With Collagen Hydrogel Solution

While it was easy to microprint antibodies, the $\mu$CP of NGF alone did not give a positive signal (Figure 2B). To establish the $\mu$CP of a protein (NGF) on the semipermeable (0.4 µm pore) membranes, we used a well-established technique to produce collagen hydrogels. While collagen alone gave the background only (Figure 2A), the $\mu$CP of the NGF loaded in the collagen hydrogel solution showed a strong and clear print and many lines after staining with an Alexa-546 anti-NGF antibody (Figure 2C). Again, no signal was seen in the green channel (Figure 2D). To optimize the NGF $\mu$CP, different concentrations of NGF were tested, and 1,000 ng NGF per load gave the best results (Figure 2E, 229 ± 8 optical density, n = 3). The $\mu$CP of 100 ng NGF (Figure 2F, 65 ± 9 optical density, n = 3) and 10 ng NGF (Figure 2G, 15 ± 5 optical density, n = 3) significantly showed a weaker signal. To demonstrate the stability of the NGF $\mu$CP, the membranes were incubated in a “slice medium” for up to 2 weeks and the stability did not change after 9 days (Figure 2I) but markedly decreased after 14 days (Figure 2J), compared with the control (Figure 2H). After 6 weeks of incubation, the NGF $\mu$CP was no longer detectable (8 ± 1 optical density, n = 3). Compared with the Isopore membranes (HTTP022500), the $\mu$CP of NGF onto the Omipore membranes (JHWP02500) and LCR PTFE membranes (FHLC02500) was 3.3x less effective (data not shown).

Characterization of Slices in Contact With $\mu$CP

The brain slices from postnatal mice were incubated on semipermeable membrane inserts with pre-prepared $\mu$CP (Figure 3A). The NGF lines were printed on semipermeable extra membranes and the brain slices were connected to the $\mu$CP (Figure 3B). Cholinergic neurons were stained for ChAT and several ChAT+ nBM neurons survived when incubated with 100 ng/ml NGF in the medium (Figure 3C) but not without NGF (Figure 3D). The brain slices were stained with the blue fluorescent nuclear dye DAPI (Figure 3E) and connected to the NGF $\mu$CP stained with an Alexa-546 antibody (Figure 3F). Figure 3G shows that the blue fluorescent slice was directly connected to the red fluorescent NGF $\mu$CP lines.

Characterization of Cholinergic Nerve Fibers Along the NGF $\mu$CP

In the next step, we characterized the ChAT+ processes growing out of the brain slices (Figure 4A). Three weeks after incubation on NGF $\mu$CP membranes, several ChAT+ cholinergic neurons were visible and several ChAT+ processes grew toward the NGF $\mu$CP (Figure 4B). The ChAT+ extensions were clearly nerve fibers (Figure 4C) that grew along the NGF $\mu$CP (Figures 4D,E). As an additional proof of specificity, Figures 4F–H show a $\mu$CP that was not homogenously printed and not intact (Figure 4G, lines b-d). The processes stopped growing on the not intact or damaged areas (Figure 4F, arrows), while the processes grew a long distance on the intact printed lines (Figure 4F, line a; and Figure 4H).

The number of processes outside the slice and along the NGF $\mu$CP was 24 ± 2 (n = 12) processes per slice and the average length of the processes was 286 ± 35 µm (n = 12) evaluated after 3 weeks in culture. The outgrowth slightly decreased after 6 weeks in culture: 16 ± 2 (n = 12; p < 0.01) processes/slice and 205 ± 19 µm (n = 12, not significant) length of fibers.

The co-localization experiments showed that the neuronal MAP-2+ immunoreactivity (Figure 5B) partly overlapped with the cholinergic ChAT+ staining (Figures 5A,C), though MAP-2 was less strongly expressed than ChAT. The overlaps were seen within the slice, but also in the ChAT+ processes (Figure 5C). An intense GFAP staining was observed in the brain slices (Figure 5E) which did not show any staining in the ChAT+ processes (Figures 5D,F). The co-staining of cholinergic ChAT+ processes (Figure 5G) with the low-affinity NGF receptor p75NTR (Figure 5H) shows strong immunoreactivity partly in the cholinergic processes (Figure 5I).

Re-organization Within the Brain Slices

For the methodological issues, the brain slices were placed directly into the NGF $\mu$CP (Figure 6A). Besides the axonal growth, we also could see a strong re-organization of cells within the brain slices. There appeared a slight re-organization of the ChAT+ neurons and fibers along the $\mu$CP within the slices (Figure 6B). The GFAP+ cells and extensions were organized along the NGF $\mu$CP in the slices, leaving some blank lines in between (Figures 6C,D). No such re-organization was seen for the microglial Iba1+ cells (data not shown) or laminin+ vessels (data not shown).

Migration of Cells Along the $\mu$CP

Besides the numerous processes outside of the slices, we also observed that cells migrated along the $\mu$CP lines. Figures 7A–C show an example wherein DAPI+ nuclei are found along a $\mu$CP (Figure 7B), co-localizing with the ChAT+ stainings (Figures 7A,C). No GFAP+ astrocytes migrated outside the brain slices (Figure 7D), but microglial Iba1+ (Figure 7E), as well as more laminin+ endothelial cells (Figure 7F), migrated along the collagen $\mu$CP.

DISCUSSION

In the present study, we show for the first time that we can microprint the protein NGF onto semipermeable membranes.
FIGURE 3 | Coupling of brain slices to microcontact prints. Chopper brain slices (300 µm) were prepared from the basal nucleus of Meynert (nBM) and cultured on semipermeable membrane inserts (A). The slices were coupled with the µCP of NGF (B). After 2 weeks in culture, the cholinergic neurons were stained for choline acetyltransferase (ChAT) and many cholinergic neurons survived when incubated with 100 ng/ml NGF in the medium (C), while no cholinergic neurons were visible when incubated without NGF (D). The brain slices were stained for nuclear 4′,6-diamidino-2-phenylindole (DAPI) (E), blue] and were connected to the NGF µCP (stained by anti-NGF Alexa-546, red), showing that the slice grows directly in connection with the µCP. Scale bar in A = 500 µm (A), 130 µm (C,D), 250 µm (E–G).
FIGURE 4 | The outgrowth of cholinergic neurons from the brain slices coupled with NGF microcontact prints (µCP). Brain slices (300 µm thick) of the nBM were prepared and connected to the NGF µCP, incubated for 1 week with 100 ng/ml of NGF in the medium, and then further for 2 weeks without NGF. The slices were fixed and stained for ChAT and Alexa-488 (green) and the ChAT processes were evaluated at the borders of the slices (A). (B) The outgrowth of ChAT+ neurons along the µCP (white lines, star *). Three typical outgrowths are seen, thin axons, but also thicker processes. (C–E) A typical example of an outgrowth of ChAT+ nerve fibers (C, green, Alexa-488), co-stained with an anti-NGF antibody (D, red, Alexa-546). (E) Shows the merged picture and that the ChAT+ nerve fibers grow along the NGF µCP. (F–H) The selectivity of growth along the µCP. (F) An example of an incomplete printing (lanes b–d), while on a complete print, (lane a) the ChAT+ processes are seen (F, arrows). Quantitative analysis shows the outgrowth of 24 ± 2 ChAT+ nerve fibers per slice with a length of 286 ± 35 µm after 3 weeks in culture. Ten brain slices were prepared from five animals. Scale bar in A = 270 µm (A), 60 µm (B), 54 µm (C–E), 108 µm (F–H).
FIGURE 5 | Co-localization of cholinergic neurons stained with ChAT (A,D,G) and with microtubule-associated protein-2 [MAP-2, (B)] or glial fibrillary acidic protein [GFAP, (E)] or the low-affinity NGF receptor p75NTR (H). Brain slices were cultured on NGF μCP for 3 weeks, fixed, and stained for ChAT (Alexa-488, green) and MAP-2, GFAP, or p75NTR (Alexa-546, red). The merged pictures (C,F,I) show the orientation of the slices, the borders (white thick dotted lines), and the suggested μCP lanes (white small dotted lines with a star *).

(A) The ChAT+ processes (six lanes) grew out of the slice (arrow) along the μCP. (B) The MAP-2+ processes which fully co-localized with the ChAT+ processes and display a differential expression intensity [see as an example the white circle in (B)]. (D) ChAT+ processes (4 lanes) growing out of the slices (arrow) along the μCP. (E) shows intense GFAP+ staining within the brain slices, more extensive at the borders representing reactive astroglisis, but nearly no immunopositive processes outside the slices [see the white circle in (E)]. (G) The immunostaining of two selected ChAT+ processes along a μCP, which fully co-localizes with the p75NTR neurotrophin receptor (H). The small arrow in (G–I) point to strong differential p75NTR immunoreactive varicosities (probably synaptic processes) within the ChAT+ processes, but also strong neuronal immunoreactive areas [white circle in (H,I)] are found. Note that all these stainings do not come from the same slice, but from four to six brain slices prepared from two to five animals. Scale bar in A = 180 μm (A–F), 36 μm (G–I).
when loaded into the biomaterial collagen. We could connect a brain slice to the NGF μCP and provide data that cholinergic processes grew along the NGF μCP and that astrocytes re-organized in the brain slices.

**Organotypic Brain Slices as a Tool to Study Axonal Growth**

Organotypic brain slice cultures represent a physiologically relevant three-dimensional *ex vivo* model [see our review (20)]. As the main architecture of the cells is preserved, it allows the *in vitro* investigation of cellular and molecular processes in the brain areas of interest. The group of Gähwiler was the first who succeeded in culturing organotypic brain slices utilizing the roller-tube technique (31). This technique was modified, maintaining organotypic brain slices cultured on semipermeable membranes (19). In our lab, we used the “chopper” technique and have extensively studied the cholinergic neurons of nBM (9, 11, 21–24, 30), dopaminergic neurons (1, 30, 32), or serotonergic neurons (33). A major benefit of organotypic brain slice cultures is the opportunity to perform co-culture experiments, which allows the study of two or more related brain areas. Dopaminergic nerve fiber growth between two co-slices has been reported by several research groups (34–36) including ours (1, 32), enabling the investigation of dopaminergic neurons concerning Parkinson’s disease. Similarly, we showed that the nerve fibers of cholinergic neurons from the nBM grow toward the cortex in organotypic brain slice co-cultures (37). Organotypic brain slice cultures were prepared from postnatal day 8–10 brains as the tissue and cell survival is high in this timeframe (37). The older the donor animals (>14 days postnatal), the more tissue and cell death occur in culture. The brains of younger donor animals exhibit a looser texture and morphology, as they are more immature. The slices were stained at the earliest after 2 weeks *in vitro*, as they need this period to completely flatten from 300 to ~100 μm (20). The grade of flattening is not only an important parameter of macroscopic cell

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**FIGURE 6** | Re-organization within the slices placed onto the NGF μCP (A). Note some form of the re-organization of ChAT+ cholinergic neurons and fibers (B) along the μCP [(B), dotted lines, star *]. The GFAP+ cells and processes re-organized in the slice and grew on the NGF μCP [(C), dotted white line] leaving some empty areas in-between (D). For each representative two to four brain slices were prepared from two to five animals. Scale bar in A = 210 μm (A), 50 μm (B,C), 25 μm (D).
survival but also increases the quality of immunostaining and microscopic analysis. The thicker the slices, the less antibodies are able to diffuse deep into the slices and the images may appear more blurry. However, it is an advantage that we performed all immunostainings free-floating, as antibodies are then able to penetrate the slice from both sides improving image quality.

**Microcontact Printing of Pure NGF Alone**

Microcontact printing allows cellular engineering by printing proteins of interest in defined stripes and it is a simple and efficient method to pattern different surfaces with a wide range of proteins. In the present study, we aimed to microprint NGF alone onto semipermeable membranes to study cholinergic nerve fiber growth in vitro. First, we reproduced a μCP of fluorescently labeled antibodies onto a semipermeable membrane to demonstrate that printing still works in our study (1). The antibodies preferentially attach to the hydrophobic PDMS stamp and are excellently transferable, thereby confirming that μCP is an efficient method for protein transfer to semipermeable membranes. The μCP of NGF alone onto semipermeable membranes, however, was not efficient. We have already observed this problem before, trying to microprint GDNF protein alone onto semipermeable membranes (1). To study the outgrowth of dopaminergic nerve fibers, we printed anti-GDNF antibodies onto the semipermeable membrane and loaded them with the GDNF protein (1). As far as we know, the microprint of pure NGF onto semipermeable membranes or any other substrate has not been published yet and the microprint...
of pure NGF alone onto semipermeable membranes was not
effective in our study. This may have several reasons, namely,
the (a) inefficient adsorption of NGF to the surface of the PDMS
stamp, (b) inefficient transfer of NGF to the semipermeable
membrane, (c) diffusion and loss of NGF through the pores of the
semipermeable membrane, (d) disruption of the NGF molecule
structure during printing procedure, or (e) blockade of the anti-
NGF antibody recognition region preventing its detection.

Collagen as a Carrier to Microprint NGF
Collagen is the main component of connective tissue, accounting
for almost one-fourth of the total body protein in humans (38),
and in the nervous system, it supports cell differentiation,
attachment, migration, proliferation, and survival (39). Therefore,
collagen is highly suitable as a raw material for tissue-
engineered scaffolds providing a biocompatible, biodegradable,
non-toxic, and versatile possibility to mimic both the structural
and biological properties. Our laboratory has already acquired
a lot of experience in working with collagen (40). So far, we
were able to load collagen hydrogels in organotypic brain slice
cultures with NGF (11), fibroblast growth factor-2 (26), and
GDNF (1) taking advantage of the PEG crosslinking system. In
contrast to our earlier study (11), we aimed not only to protect
the cholinergic neurons in the nBM via NGF application but also
to initiate axonal growth in defined directions. Therefore, we
microprinted a collagen hydrogel solution loaded with NGF to
immobilize NGF in the pattern next to the slice enabling guided
neural fiber outgrowth. In the present study, we succeeded in
microprinting NGF loaded in collagen onto 0.4 µm pore
semipermeable membranes. To our knowledge, collagen has
never been microprinted together with any other protein onto
semipermeable membranes. There are two reports showing
techniques in applying NGF in combination with collagen,
however, it was “molecular printed” to a collagen gel (41)
and “3D bioprinted” (42). In our present study, we verified
specificity via immunostainings for NGF with fluorescent
antibodies. Furthermore, the quantitative evaluation showed
that the µCP of NGF was dependent on the amount of NGF
loaded. Thus, the microprinting of NGF onto semipermeable
membranes using collagen hydrogel solution was shown by
this study for the first time. This method is easy, fast, cheap,
and versatile as nearly any protein can be printed, which is
shown by our preliminary experiments, e.g., we can microprint
a 50 kDa tau protein but also a 4 kDa β-amyloid peptide
(Supplementary Figure 1).

Cholinergic Neurons and NGF-Induced
Axonal Growth Along NGF µCP
Cholinergic neurons are located in the nBM and the medial
septum providing projections to the entire neocortex and
hippocampus, respectively (4, 5). The neurotransmitter
acetylcholine has a relevant impact on memory and a loss of
acetylcholine in AD directly correlates to memory loss (6, 7).
Cholinergic neurons were immunohistochemically stained
for the enzyme ChAT as this enzyme is exclusively expressed
in the cytosol of cholinergic neurons and we have extensive
experience with such stainings (21, 37). Cholinergic neurons
express the neurotrophin receptor p75NTR which almost entirely
co-localizes with ChAT in the nBM, as well as the medial septum
(43). The NGF is a target-derived neurotrophic factor and it is
well-known that NGF is essential for the viability of cholinergic
neurons (9–11). In the present study, we confirmed previous
work and showed that the µCP of NGF supports the survival and
subsequent nerve fiber growth of nBM neurons in organotypic
brain slices.

Commonly, the extracellular matrix proteins poly-lysine,
fibronectin, and laminin are microprinted onto glass slides or
Petri dishes before neuronal cell populations are seeded (44–46).
As those proteins are known to support neural cell growth and
differentiation, this approach is frequently used to study neurons
in cell culture. Taking advantage of the same approach, others
succeeded in µCP semaphorins (47), ephrins (48), and netrins
(49) onto glass slides or plastic dishes. All of them are molecules
known to effectively direct axon guidance. In the present study,
we used the target-derived neurotrophic factor NGF to guide
cholinergic nerve fibers along the µCP lines.

To connect the organotypic brain slices to the µCP, freshly
prepared slices were carefully placed onto the µCP membrane.
The slices were positioned in the center of the µCP, which
enabled us to investigate neuronal growth in two directions
per slice. The most difficult point was to ensure that the slices
had obtained the desired orientation on the µCP, as the µCP
itself was hardly or not visible at all during the arranging
procedure. The marks on the membrane assisted us in assuming
the localization of the µCP and we were able to positively connect
the slices and µCP as evidenced by the DAPI-stained cells of a
slice superimposing a µCP. To test the neuronal growth of
cholinergic neurons, a brain slice containing a region of nBM
was placed onto a NGF µCP. Our analysis clearly showed
selective ChAT+ outgrowths from the slice along the NGF
printed regions. Their appearance, however, was very diverse,
ranging from single nerve fibers and nerve fiber bundles to
thick nerve fiber networks. The diversity of outgrowths may
result from the composition of the slice itself, depending on
which cells and factors are more highly enriched along the
NGF µCP at the beginning of the culture. We found not only
outgrowths but also a ChAT+ fiber reorganization along the
NGF µCP within the brain slice, pointing toward a highly
adaptable system. Moreover, we observed ChAT+ cells and,
occasionally, an area of cells with unknown origins which
migrated along the NGF µCP as well. This underscores the
importance of NGF for cholinergic neurons, but also that
NGF appears to affect other neuronal cell mechanisms as well.
The neuronal marker MAP-2 co-localized with the ChAT+
processes, which strengthens the assumption of neuronal growth
along NGF µCP. Additional immunohistochemical staining
using p75NTR antibodies showed strong colocalization with
ChAT+ processes providing additional evidence that cholinergic
neurons grew along the NGF µCP. No co-staining was seen
for astroglial GFAP, microglial Iba-1, and vascular laminin. It
is likely that the collagen per se provides a potent substrate
to stimulate growth along the microprints. The ChAT+ nerve
fibers grew \( \sim280\mu\text{m} \) out of the brain slices after 3 weeks but did not extend further after 6 weeks. It was assumed that the collagen degrades completely within 14 days, but the outgrowing processes survive on the membrane for longer periods. No more growth was seen after 3 weeks, probably due to the lack of NGF microprints. To our knowledge, our study is the first to observe neuronal growth along NGF \( \mu \text{CP} \) in slices.

**Intra-Slice Re-organization on NGF Microcontact Prints**

For technical reasons, the slices were positioned at the center of the \( \mu \text{CP} \) as this was easier and guaranteed that the slices were clearly connected to the \( \mu \text{CP} \). Thus, this strategy enabled us not only to visualize outgrowth but also to study any internal reorganization within the slice along the NGF \( \mu \text{CP} \).

The most interesting observation was the strong astrocytic (GFAP+) internal reorganization along the NGF \( \mu \text{CP} \) after 3 weeks in culture. Furthermore, our analysis showed that in some parts, cholinergic (ChAT+) internal reorganization took place along the NGF \( \mu \text{CP} \). In contrast, no vessel (Laminin+) and microglial (Iba-1+) internal reorganization were visible. The reasons for the NGF-mediated re-organization of astrocytes are entirely ambiguous. There is evidence that NGF is produced and released by astrocytes after inflammation, suggesting an autocrine or paracrine mechanism (50–52). Further, NGF induces the expression of the \( p75^{NTR} \) receptor in astrocytes especially during development, inflammation, and after injury (50, 53). More importantly, it has been reported that NGF facilitates astrocytic migration via the \( p75^{NTR} \) as the trkA receptor is not expressed in astrocytes (53, 54). In fact, NGF can also increase the migration of multipotent astrocytic stem cells (55), but also possibly of oligodendroglia expressing \( p75^{NTR} \) and associated with radially oriented astroglia (56). The process of astrocytic migration is unclear, but it seems to be particularly essential for the structural organization during development or scar formation after brain damage (57). Our data provide evidence of the self-reorganization of astrocytes along the NGF \( \mu \text{CP} \), considering that NGF may be a strong chemoattracting growth factor for astrocytes. However, we cannot exclude that collagen alone or in combination with NGF is responsible for the self-reorganization of the astrocytes. It has been seen that astrocytes interacted and aligned with the \( \mu \text{CP} \) of laminin (58). The self-reorganization of the astrocytes may also force cholinergic (ChAT+) cells to migrate and organize along the NGF \( \mu \text{CP} \) (or the opposite way round).

**Migration of Cells Along the \( \mu \text{CP} \) Lanes**

In our study, we observed some migration of cells. It seems possible that cholinergic ChAT+ neurons migrated along the \( \mu \text{CP}(\text{NGF}) \) lanes. It seems very likely that NGF supports some form of cellular migration, although this mechanism is not fully understood. Especially, it is novel to see the migration of fully differentiated postnatal cholinergic neurons. We further showed that endothelial cells or microglia migrated along the collagen \( \mu \text{CP} \). On the other hand, we can exclude migration of astroglia, as we did not see any staining along the \( \mu \text{CP} \) lanes. Regarding microglia and endothelial cells, we do not suggest an NGF-dependent process. Rather, it is possible that the carrier material collagen strongly supports the differentiation, attachment, migration, proliferation, and survival of those cells (39). In fact, it has been well-documented that extracellular matrix proteins, such as poly-lysine, fibronectin, and laminin, appear to initiate similar behavior in different cell types (44–46). Astrocytes do not migrate or extend processes along the \( \mu \text{CP} \), which could be explained by the (a) downregulated GFAP expression in those areas, (b) different subtypes of astrocytes that do not express GFAP, or (c) a switch from reactive to inactive state and degradation of GFAP. Definitely, GFAP represents the reactive astrocytes within the slices, which are more pronounced at the borders. Alternatively, GFAP+ cells may need a longer time in culture to migrate the same distance as other cells (45). More work is necessary to study the migratory capacity along the \( \mu \text{CP} \).

**Limits of the Study and Outlook**

This work has some limits: (a) Collagen is a potent biomaterial, but we observed some minor changes in the composition of the hydrogels, which could influence the results. Collagen degrades over time and releases NGF within 2 weeks, thus it seems likely that the outgrowth along NGF \( \mu \text{CP} \) occurred within 2 weeks. (b) In our study, we have not performed retrograde tracing experiments. So, it would be interesting to apply a fluorescent dye and to observe if it is retrogradely transported to the slices, such a dye could be MiniRuby. (c) In this respect, the live-cell imaging of such tracing could also be of interest to observe the retrograde transport of the fluorescent dye. (d) Although we found a relatively good outgrowth of \( \sim300\mu\text{m} \) over 3–6 weeks, it is not clear if the outgrowth is time-dependent and can be extended. Partly it was hard to differentiate the border of the brain slices and to measure the distance length. (e) In our experiments we used NGF in the medium to prime the initial survival of cholinergic neurons; we have not tested if the survival on \( \mu \text{CP}(\text{NGF}) \) alone is sufficient. In addition, the dissection of the nBM brain area was not easy and it can be possible to collect and culture slices that do not contain cholinergic neurons. To enhance the chance, two brain slices were placed on a membrane. The quality of the slices was always visualized optically and a good flattening and high transparency are an indicator of a good slice preparation. Thicker slices and slices without any cholinergic neurons were deleted from this study.

The method of \( \mu \text{CP} \) in combination with organotypic brain slices is very potent and promising and has several future applications: (a) First, principally we can microprint any protein or peptide of interest (in preliminary experiments we microprinted tau and A\(\beta\), Supplementary Figure 1). This increases the potential of the method markedly, as any commercial protein or peptide can be loaded. (b) Such a broad range may also allow us to microprint a cocktail of different proteins or peptides at the same time. (c) The method of \( \mu \text{CP} \) may also allow the printing of different patterns in different directions, which again markedly enhances the possibilities...
to study neuronal fiber growth. (d) Although not tested, we may also consider printing cells, such as isolated astrocytes or stem cells. (e) This method of µCP may also allow us to study vessels and their arborization out of the brain slices and the possible re-connection of vessels. (f) This method of µCP may have the potential to induce the migration of microglial cells along a printed line with small 1.0 µm beads which can be incorporated. (g) The use of different biomaterials may potentiate the µCP pattern and may further improve the method. (h) Such a method may be useful to study the neuronal re-growth of cholinergic neurons in AD, but also of dopaminergic neurons in Parkinson’s disease. Alternatively, the re-growth and connection of spinal cord nerve fibers may also be studied. (i) Finally, this may markedly reduce animal experiments as many slices (depending on thickness and brain region of interest) can be prepared from one brain contributing to the 3Rs.

In summary, we showed for the first time that we can microprint NGF onto 0.4 µm semipermeable membranes using collagen as a loading vehicle. The cholinergic nBM organotypic brain slices survived and the cholinergic nerve fibers extended toward the NGF µCP. We also showed that GFAP+ astrocytes re-organized in the slices on the NGF µCP. We showed here for the first time that we can link µCP with brain slices and this method has a high capability to study different aspects in neurobiology. As this technique is easy, fast, and cheap, we can print nearly any protein/peptide onto a membrane. This technique will allow studying nerve fiber growth but may also be suitable to study vessel growth or migration of cells.

**DATA AVAILABILITY STATEMENT**

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

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**ETHICS STATEMENT**

Ethical review and approval was not required for the animal study because all experiments conformed to Austrian guidelines on the ethical use of animals and were in line with the 3Rs rule as all efforts were made to reduce the number of animals. All animal experiments were defined as Organentnahme according to Austrian laws.

**AUTHOR CONTRIBUTIONS**

KS performed all experiments and wrote the manuscript. CH designed and analyzed the data and wrote the manuscript. Both authors contributed to the article and approved the submitted version.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fneur.2021.775621/full#supplementary-material
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