Neuromuscular Contacts Induce Nitric Oxide Signals in Skeletal Myotubes in vitro

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Abstract
It has previously been shown that skeletal myotubes express nitric oxide synthase (NOS) and produce and release NO signals. NOS is also part of agrin-induced acetylcholine receptor aggregations on myotubes. As nerve-muscle interactions underlie reciprocal signaling mechanisms, we hypothesized that NO signals in target myotubes may be induced by neuromuscular contacts in development. Chimeric neuron-myotube co-cultures were prepared using p75-selected spinal cord neurons from embryonic chicken. Confocal imaging revealed robust 1,2-diaminoanthraquinone red fluorescence indicative of de novo formation of NO only in those myotubes which were contacted by neurites, also verified by pre- and postsynaptic marker costaining (anti-synaptotagmin and α-bungarotoxin). Neither soluble agrin nor sensory dorsal root ganglionic neurons showed comparable effects in this model. We concluded that in target skeletal muscle cells the NOS/NO system is controlled by motoneuron contacts by as yet incompletely understood signaling mechanisms. Endogenous NO signaling in myotubes may be essential during synapse formation and plasticity of the neuromuscular system.

Introduction
Nitric oxide (NO) signals are fundamental in the development and function of the central and peripheral nervous system [1]. The dimeric and gaseous NO molecules produced by NO synthases (NOS) act as freely diffusible messengers in defined neurons in the central and peripheral nervous system [2]. NO addresses cellular and subcellular targets in defined neuronal networks [3, 4], e.g. nitrosylation [5]. The NOS gene has a complex structure [6] and three major isoforms of NOS, neuronal (NOS-1), macrophage (NOS-2) and endothelial (NOS-3), have been cloned and characterized [7]. NOS is abundantly expressed in peripherally innervated organs suggesting important functions of target-derived NO signals [8].

While somatic motoneurons in the spinal cord normally lack NOS expression [9] NOS is abundantly found in their target skeletal muscle fibers representing a major
physiological source of NO in the body [10–13]. In the muscle fiber, NO is generated by Ca$^{2+}$-dependent constitutively expressed NOS-1 or by one of its coexpressed muscle-specific splice variants termed NOS-μ [14]; both are associated with the sarcolemmal dystrophin-associated glycoprotein complex (DAGC) of normal adult muscle fibers [15]. NOS-1 is concentrated at the neuromuscular junction [16–18] together with cyclic GMP-dependent protein kinases and soluble guanylyl cyclase [19, 20] and is part of agrin-induced nicotinic acetylcholine receptor (AChR) aggregations on myotubes including NMDA-receptor, ion channels, PSD-95 or 43K-rapsyn linker proteins, suggesting fundamental signaling mechanisms of NO in neuromuscular synapse formation [21–24].

Skeletal mouse C2C12 myotubes express a concomitant cytosolic pattern of NOS-1–NOS-3 isoforms in culture [25]. During postnatal development, NOS becomes redistributed during functional skeletal muscle innervation [26]. Endogenous NO is a retrograde signal involved in nerve terminal differentiation and neuromuscular junction plasticity [27, 28] and development [29]. Whether endogenous formation of NO molecules in target skeletal myotubes is controlled by neurons is presently unknown. We, therefore, prepared chimeric neuron/myotube co-cultures in order to determine whether NO formation in skeletal myotubes is induced by neuronal contacts. Viable C2C12 myotubes were preloaded with 1,2-diaminoanthraquinone (DAQ), a red-fluorescent NO-specific probe for intracellular detection. The DAQ-preloaded myotubes were co-cultured with embryonic chicken p75-selected spinal cord motoneurons or sensory dorsal root ganglionic (DRG) neurons. In p75 neuron/myotube co-cultures, robust DAQ red fluorescence staining was detectable in the sarcosol of target myotubes by confocal laser microscopy thus indicating intracellular de novo formation of NO. Soluble agrin or sensory neurons failed to show similar effects in this model. Our in vitro experiments confirmed that endogenous NO synthesis in target myotubes might be under the control of neuromuscular contacts of somatic motoneurons.

**Materials and Methods**

Mouse skeletal C2C12 myoblasts (ATTC, Rockville, Md., USA) were cultured in proliferation medium [DMEM with 4.5 g/l glucose/1% non-essential amino acids (Biochrom, Berlin, Germany)/1% penicillin-streptomycin, P/S (Gibco, Karlsruhe, Germany)/1% L-glutamine (Biochrom)/10% fetal calf serum (FCS; PAA Laboratories, Marburg, Germany) under 6% CO$_2$ atmosphere in a 37°C incubator (humidity 95%)]. Myotube formation was induced within 2–3 days in vitro following exchange of the fusion medium (DMEM 1 g/l glucose/1% L-glutamine/1% P/S/2% FCS), as previously described [23].

Five- to 6-day-old chick embryos (E5/E6) were obtained from fertilized white leghorn eggs and dissected under sterile conditions. Embryonic trunks were dissected at the ventral midline, the thoraco-lumbar spinal cord enlargements Th12 to L5 were removed following laminectomy and discretely microdissected into ventral horn tissue columns incubated in ice-cold Hank’s salt solution (HSS, Gibco), washed twice with ice-cold HSS, and incubated in 0.025% trypsin/HSS (37°C water bath for 30 min). The tissue pieces were dissociated and triturated in 1 ml of DMEM/2% FCS/DNase I (0.5 mg/ml, Boehringer, Mannheim, Germany) by siliconized glass pipettes. In addition, lumbar (L1–L5) DRG neurons were isolated from E8 chicken embryos. Chick embryonic neurons were purified by preplating steps as previously described [30].

A rabbit anti-mouse antibody which recognizes the extracellular domain of the p75 low-affinity nerve growth factor receptor (NGFR) expressed by the pool of spinal motoneuron cells at developmental stages E5/E6 [31] and which is cross-reactive with the chicken p75 NGFR (Chemicon, Hofheim, Germany) was used for ‘cell panning’: 35-mm culture dishes (Nunc, Roskilde, Denmark) were first coated with a rabbit-IgG Fc-specific antibody (Dianova, Hamburg, Germany) dissolved in PBS (1:150) for 2 h at room temperature (RT). Following three washes in PBS, the dishes were incubated with rabbit anti-mouse p75 NGFR antibody, diluted 1:100 in PBS for 2 h at RT. After three washing steps with DMEM/2% FCS, 2 ml of the pre-plated neuron-like cells were seeded into p75 NGFR-precoated 35-mm dishes (seeding rate: 200,000 cells/dish) and incubated for 30 min at RT. The neuronal cells captured by immobilized p75 NGFR antibodies at the bottom of the dish were mechanically remobilized and harvested in 2 ml of DMEM/2% FCS, centrifuged (300 g, 2 min) and recounted. Cell panning yielded about 250,000 living cells/ml (Trypan blue exclusion test), about 92.2 ± 1.2% (n = 6, two independent experiments) of these cells were immunopositive for the E5/E6 chick motoneuron-specific marker calcitonin-gene related peptide (anti-CGRP monoclonal antibody, 1:2,500 in PBS, Amersham, Amersham, UK) as described [32].

Purified neurons were seeded onto fused C2C12 myotube monolayers grown onto uncoated glass coverslips (seeding rate: 1,000 neurons/13-mm-diameter coverslip) and cocultivated in DMEM/2% FCS supplemented with 2% chicken embryo extract (Gibco) and raised as chick-mouse chimeric co-cultures for up to 8 days in vitro. Neurites were immunostained with anti-neurofilament (NF) H polyclonal antibodies (Chemicon), diluted 1:500 in PBS, for 2 h at RT followed by secondary Cy2-conjugated anti-rabbit antibody (Dianova) diluted 1:100 for 1 h at RT. As presynaptic marker, we used a monoclonal antibody directed against the 65-kDa band of the synaptic vesicle protein synaptotagmin, which is identical in humans, rats and chicken (clone ASV 48, StressGen, Victoria, Canada), diluted 1:200 in PBS, for 2 h at RT. After three washing steps with DMEM/2% FCS, 2 ml of the pre-plated neuron-like cells were seeded into p75 NGFR-precoated 35-mm dishes (seeding rate: 200,000 cells/dish) and incubated for 30 min at RT. The neuronal cells captured by immobilized p75 NGFR antibodies at the bottom of the dish were mechanically remobilized and harvested in 2 ml of DMEM/2% FCS, centrifuged (300 g, 2 min) and recounted. Cell panning yielded about 250,000 living cells/ml (Trypan blue exclusion test), about 92.2 ± 1.2% (n = 6, two independent experiments) of these cells were immunopositive for the E5/E6 chick motoneuron-specific marker calcitonin-gene related peptide (anti-CGRP monoclonal antibody, 1:2,500 in PBS, Amersham, Amersham, UK) as described [32].

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10 μm myotube segments (n = 40) in (i) motoneuron-myotube co-cultures, (ii) aneural myotubes and (iii) myotube cultures supplemented overnight with chick motoneuron conditioned medium. Values are given as means ± SEM.

NO formation was measured in supernatant culture medium using a quantitative colorimetric NO detection kit (Cayman Chemical, Ann Arbor, Mich., USA) based on the photometric nitrate/nitrite assay with the Griess reagent (detection limit 2 μM) in 96-well titer plates (Nunc) according to the manufacturer’s protocol. NO formation was inhibited in fused myotubes cultured for 2–3 days in vitro in plates (Nunc) according to the manufacturer’s protocol. NO formation was inhibited in fused myotubes cultured for 2–3 days in vitro. NO formation was inhibited in fused myotubes cultured for 2–3 days in vitro. NO formation was inhibited in fused myotubes cultured for 2–3 days in vitro. NO formation was inhibited in fused myotubes cultured for 2–3 days in vitro. NO formation was inhibited in fused myotubes cultured for 2–3 days in vitro. NO formation was inhibited in fused myotubes cultured for 2–3 days in vitro. NO formation was inhibited in fused myotubes cultured for 2–3 days in vitro.

Results

Survival and neurite outgrowth of p75-selected spinal cord motoneurons were evident 48–72 h after seeding as verified by synaptotagmin-immunopositive neural somata and neuritic processes expanding over elongated myotubes (fig. 1A). Double immunolabeling showed a selective clustering of AChR on the myotube surface verified by postsynaptic marker αBGTx at sites of neurite contacts verified by the presynaptic immunomarker synaptotagmin (fig. 1B). At higher magnifications of cluster areas, sequential image analysis confirmed the close spatial relationship between outgrowing neurites and AChR aggregates on myotubes (fig. 1C, D). The yellow pixels generated by region-oriented segmentation image analysis reflect a 1:1 overlap of fluorescent-labeled structures of the neurite process and of AChR aggregates at sites of intimate contacts (fig. 1D).

The overall pattern and amount of AChR aggregates on myotubes from neuron-myotube co-cultures (fig. 2A) are very different from aneural myotubes (fig. 2B), and from those myotubes which were grown in the presence of p75 neuron-conditioned medium. The quantitative evaluation showed an increase in the number of specific AChR clusters (i.e. macroclusters) on defined myotube segments (3.9 ± 0.3 clusters/100-μm segment, n = 60) in co-cultures (fig. 2C) when compared to aneural cultures, i.e. myotubes cultured under identical conditions without neurons (0.6 ± 0.3 clusters/segment, n = 40). Addition of neuron-conditioned medium containing soluble factors released by neurons did not result in comparable membrane clustering (fig. 2C).

In order to investigate the time course of early neuron/myotube interactions, the degree of specific BGTx cluster patterns on myotubes was counted. Co-clustering was detected starting from day 5 after seeding (fig. 2D). A significant increase in the amount of BGTx-cluster-positive myotubes was however detected 5–6 days in vitro after seeding of p75 motoneurons reflecting onset of major triggering mechanisms supporting AChR membrane clustering. As neurites grew out in co-cultures as early as 48 h...
after seeding (anti-NF immunomarker), we concluded that neurite-myotube contacts must have established after 3–5 days in vitro. A decline in the BGTx clusters was noted in co-cultures at later developmental stages in vitro.

We next blocked NO biosynthesis in myotube cultures by treatment of fused myotubes with various concentrations of known NOS inhibitors followed by colorimetric NO assay (fig. 3). Blocking with variable final concentrations depending on NOS inhibitors decreased the concentration of NO in myotube cultures as compared to myotube-only cultures without blocking treatments (n = 6). In the absence of NOS blockers, NO levels were significantly elevated in fused myotubes and further increased in co-cultures (fig. 3).

The formation of intracellular NO was directly visualized in individual myotubes by preloading them with the non-fluorescent NO-sensitive compound DAQ, which, in the presence of NO, generates a strong red fluorescence detected by confocal microscopy. Quantification was done by ROI pixel intensity measurements (scale: 0–255 arbitrary units). In myotubes pretreated with the NOS inhibitor AMT, DAQ red fluorescence (verifying intracellular NO) was not detectable in the myotube sarcosol (12.25 ± 7) as determined by ROI-pixel intensity measurements, suggesting that the in vitro DAQ protocol specifically stained de novo formation of intracellular NO (fig. 4A). Without blocking, aneural myotubes revealed only weak DAQ red fluorescence (38.9 ± 9) thus reflecting intracellular basal levels of NO (fig. 4B). Robust DAQ red fluorescence (ROI pixel intensity 98.0 ± 3) was however detectable in many myotubes in co-culture thus reflecting an approximately 159% increase of the relative fluorescence at the ROIs measured compared to basal values found in other myotubes of the same co-cultures (fig. 4C). The DAQ fluorescence in responding myotubes was always diffusely distributed and, for example, not restricted to membrane aggregations possibly induced at sites of neurite-myotube contacts. Neither the addition of the AChR aggregating molecule s-agrin nor co-cultures with sensory DRG neurons showed comparable effects on myotube DAQ fluorescence (not shown).

Costaining experiments with anti-NF and DAQ verified the presence of neurite processes on strongly red fluorescent, i.e. DAQ-positive, myotubes only (fig. 4D–E, see also 4C). Neurite processes were not detectable on DAQ-negative myotubes adjacent to DAQ-positive myotubes (fig. 4E, inset). During axonal outgrowth and guidance,
Fig. 2. Epifluorescence microscopy of αBGTx-labeled specific AChR ‘macroclusters’ (white spots) determined in co-cultures (A) versus aneural myotubes with few spontaneous ‘microclusters’ (B). C The number of BGTx clusters per myotube segment in co-culture is significantly different from aneural cultures or from cultures supplemented with motoneuron-conditioned medium. D Graph shows percent changes in myotubes with BGTx-labeled AChR clusters versus cluster-positive aneural myotubes (set as 0%) after 0–8 days in co-culture. Specific AChR clustering responses are lacking on myotubes at 1–4 days in vitro (i.e. <5%) suggesting that physical contacts between neurites and myotubes were not yet established in co-culture. At 5–6 days in vitro, the number of specific BGTx-cluster-positive myotubes is significantly increased, thus monitoring neuron–myotube contacting in vitro. Bar = 40 μm.
Fig. 3. Colorimetric NO assay of the culture medium after various treatments. Minimal amounts of NO were detected in myotube cultures following administration of various NOS inhibitor concentrations: 1 μM diphénylhydroiodonium chloride (DPI), 100 nM 3-brom-o-7-nitroindazole (3-B-7-NI), 20 nM AMT or 1 μM n-o-nitro-L-arginine (L-NNA) in culture medium (5 days in vitro). Elevated NO levels were detected in co-cultures (41 μM) as compared to basal NO concentrations in myotube-only cultures (20 μM) of the same developmental stage in vitro (5 days in vitro).

Discussion

The major findings of the present study were as follows: (i) p75 neurons isolated from embryonic chick spinal cords survived and extended neurites when co-cultured with C2C12 myotubes, (ii) increased NO concentrations were determined in neuron-myotube co-cultures by colorimetric NO assay, (iii) myotubes in co-culture showed strong sarcosolic DAQ red-fluorescent stain when monitored by confocal laser microscopy, (iv) the DAQ-positive myotubes in co-culture were always contacted by neurites identified by pre- and postsynaptic markers and (v) neither agrin nor sensory neurons produced similar effects on cultured myotubes. We suggest that endogenous NO formation in myotubes is dependent on neuronal signals resulting in NOS induction at the developing neuromuscular junction. The signaling cascades responsible for myotube NOS induction remain to be clarified.

To our knowledge, this is the first report on neuronal contact-induced NO formation in skeletal myotubes visualized by the NO-sensitive fluorescent marker DAQ by confocal microscopy. Cell markers like DAQ and DAF-2 (4,5-diaminofluorescein) are useful for specific and semiquantitative visualization of de novo formation of NO in viable cells in vitro and in vivo [35, 37, 38]. Induction of NO in myotubes was preferentially detectable in co-cultures with p75-selected neurons enriched from the motoneuron pool of the embryonic chicken lumbar spinal cord. Intimate neurite-myotube contacts were identified in co-culture by the presence of postsynaptic AChR clusters on myotubes [39–43] and redistribution of ACh receptor and esterase mRNA [44–46].

Myotubes form postsynaptic myopodia (microprocesses) at sites of axonal contacts in Drosophila melanogaster embryos in vivo [47] and in neuron-myotube co-cultures within the first 24 h in vitro [43] suggesting dynamic interactions following establishment of contacts between myotube and axon in vitro and in vivo. Initial events in neuromuscular synapse formation [48, 49] include agrin-induced aggregation of membrane complexes or macroclusters including the AChR, ion channel pro-
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Fig. 4. Intracellular detection of NO in viable C2 myotubes preloaded with the NO-sensitive compound DAQ. A Preincubation with the selective NOS inhibitor AMT resulted in negative DAQ fluorescence of myotubes (MT); the myonuclei (x) are spared. Broken lines denote the location of a DAQ-negative myotube. B Aneural myotube cultures (5 days in vitro) show moderate DAQ fluorescence signals in sarcosol and myonuclei reflecting NO baseline production. Co-cultures (4–5 days in vitro) with robust DAQ fluorescence signals (red) in myotubes reflecting elevated NO production; the myonuclei are masked by the large amounts of intracellular DAQ signals. D DAQ-positive myotube with a traversing anti-αNF-immunopositive neurite (green, vertical arrows) of an adjacent p75-selected neuronal soma (horizontal arrow). E DAQ-positive myotube with Cy2 αNF-immunostained neuritic process (arrows). DAQ-positive mononucleated myoblasts (x) located next to multinucleated myotubes (MT). Inset shows a neighboring myotube with weak DAQ fluorescence (broken lines) lacking NF-immunostained neurites (green) from the same co-culture. A–E Bar = 20 μm (inset, 40 μm).

Agrin may act as one of several presynaptic signals for in vitro AChR clustering in cultured myotubes [54, 55]. At the postsynaptic membrane, NO acts as a downstream mediator of the initial agrin-induced AChR clustering process [22, 56], supporting the notion that the agrin/NO-signaling pathway is likely involved in redistribution of NOS-1 from cytosol to synaptic membrane clusters previously proposed [23].

The typical AChR macroclusters formed in vitro are known to be initiated by nerve-derived soluble agrin secretion at sites near nerve-muscle contacts [57, 58], whereas spontaneous clusters (i.e. microclusters) on myotubes may be induced by normal synthesis of muscle proteoglycans [59], laminins [60] or by muscle-derived agrin for example [61, 62]. In our experimental model, soluble agrin did not induce strong DAQ red fluorescence in myotubes. Addition of agrin to myotube cultures may not be a...
localized synaptic signal. Thus, lack of sarcosolic NO signals in agrin-treated myotubes may be due to more localized actions of agrin activity requiring parallel-operating presynaptic or postsynaptic signaling molecules from motor axons or target muscle cells for example. Myotubes with spontaneous AChR aggregates were also seen in our co-cultures, which, however, always lacked strong DAQ fluorescence and pre- and postsynaptic markers. The presence of diffusible trophic signals in the culture medium would hardly explain the cluster patterns of innervated DAQ-positive myotubes in co-culture since motorneuron-conditioned medium had no significant effects on single myotube cultures. The BGTx macroclusters induced by first-formed motor axon contacts on target myotubes on days 3–5 in vitro (i.e. agrin dependent) may be reversible at later in vitro development (i.e. becoming agrin-independent spontaneous microclusters), possibly due to the lack of appropriate molecular signaling or limited trophic support and thus incomplete local organization of the synaptic machinery, or simply by reversible neurite-contacting mechanisms at former target myotubes in our experimental co-culture system.

Synaptic actions by NO have previously been discussed in terms of retrograde muscle-to-nerve NO signaling [21]. NOS is concentrated in neuromuscular junctions of adult and developing rats, mice or humans [16–18]. Cyclic GMP-dependent protein kinase is coexpressed with NOS at the neuromuscular junction [19], and NOS-1 is either linked directly via PDZ-PDZ domains or indirectly via PSD-95 to the membrane N-methyl-D-aspartic acid receptor in the brain [63] supporting the notion that local Ca2+ influx could be the trigger for NOS activity at the synaptic multimolecular complex. More recently, the molecular association between synapsin and NOS has been characterized [64]. The presence of muscle-specific receptor tyrosine kinases, muscle-specific kinase (MuSK) [65], found in embryonic muscle and neuromuscular junctions provide possible molecular cues for transmembrane signaling processes and second messenger pathways initiated by motoneuron-muscle contacts.

As principally all major NOS isoforms were found in skeletal myoblasts, the question arises which of the NOS isoforms might be responsible for the robust and diffuse myotube NO signals detected in our co-cultures. Attempts to further characterize a ‘developmental-type’ isoform of NOS in fused C2 myotubes by NOS-isoform-specific antisense mRNA transfection experiments (NOS-1–NOS-3) failed to show clear correlations between NOS isoform expression and the known biological effects (i.e. fusion competence) observed in vitro [unpubl. observation]. Redundantly or residually expressed active NOS isoforms in early development may thus be responsible for critical
NO levels as a prerequisite for myoblast fusion and differentiation [66]. On the other hand, NOS isoform activity and thus NO levels may be regulated later in development subserving more spatial actions of NO addressing molecular targets of intracellular compartments of more differentiated skeletal muscle. As the mature pattern of skeletal muscle innervation, for example, is achieved at peri- and postnatal stages [67], it seems likely that the known redistribution of the cytosolic NOS to sarcolemma sites previously reported [25, 68] partly reflects myofiber innervation patterning in postnatal skeletal muscle differentiation.

However, the functional consequences of neurite-induced NO production in myotubes are manifold. In skeletal muscle, NO mediates excitation-contraction coupling, mitochondrial and glucose metabolism, and autoregulation of the microvascular network [13]. Freely diffusible NO reacts with several classes of proteins, soluble guanylyl cyclases, sarcoplasmic ATPases, respiratory chain complexes as well as radical oxygen species termed ROS [69, 70]. NO inhibits or activates the sarcoplasmic reticulum ryanodine receptor in muscle thereby modulating intracellular calcium levels [71]. In adult muscle from NOS-3-deficient mice, NO production and contractile functions were not impaired suggesting that NOS-1 is the major isoform which mediates intracellular NO signaling in mature muscle fibers [72]. Electrical stimulation of adult skeletal muscle preparations induces NOS-1, thus activity-dependent mechanisms may act as physiological triggers for skeletal muscle NO formation [73]. During development, NO is thought to participate in activity-dependent synaptic suppression in neuromuscular [27] or central synapses [74], in neuronal proliferation and survival in cultured neuronal cells [75] and in *Xenopus* brain [76], supporting the idea that similar events are regulated by retrograde muscle-to-nerve NO signals from skeletal muscle targets [77] critical for the assembly and maintenance of the postsynaptic apparatus [78]. In synaptogenesis and neurite outgrowth, NO acts by itself or in concert with other growth factor peptides [79]. Muscle-fiber-derived NO signals may act as intrinsic molecular mediators for postsynaptic AChR expression [80] or plasticity control in muscle differentiation and innervation patterning as recently proposed for AChR aggregations formed at prospective synaptic regions of muscle fibers [81]. In general, neurite-induced NO formation in vitro may be part of synergistic effects of muscle- and/or nerve-derived peptides, for example, acting on the cellular and molecular level at early or late synaptogenesis [82–84].

In conclusion, the present findings provide evidence for elevated NO levels in neuron-myotube co-cultures. De novo formation of endogenous NO has been directly visualized in viable target myotubes which were contacted by motor axons. Both agrin and sensory neurons failed to show comparable results in our model. Elucidation of the interactive pre- and postsynaptic molecular signals involved in NO biosynthesis during nerve-muscle interactions will help to further understand target-dependent NO signaling in development, differentiation and plasticity of the neuromuscular system.

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