Reduction of β-Amyloid Levels by Novel Protein Kinase Cε Activators

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Isomorph-specific protein kinase C (PKC) activators may be useful as therapeutic agents for the treatment of Alzheimer disease. Three new ε-specific PKC activators, made by cyclopropagation of polyunsaturated fatty acids, have been developed. These activators, AA-CP4, EPA-CP5, and DHA-CP6, activate PKCe in a dose-dependent manner. Unlike PKC activators that bind to the 1,2-diacylglycerol-binding site, such as bryostatin and phorbol esters, which produce prolonged down-regulation, the new activators produced sustained activation of PKC. When applied to cells expressing human APPSwe/PS1Δ6, which produce large quantities of β-amyloid peptide (Aβ), DCP-LA and DHA-CP6 reduced the intracellular and secreted levels of Aβ by 60–70%. In contrast to the marked activation of α-secretase produced by PKC activators in fibroblasts, the PKC activators produced only a moderate and transient activation of α-secretase in neuronal cells. However, they activated endothelin-converting enzyme to 180% of control levels, suggesting that the Aβ-lowering ability of these PKCe activators is caused by increasing the rate of Aβ degradation by endothelin-converting enzyme and not by activating nonamyloidogenic amyloid precursor protein metabolism.

Alzheimer disease is characterized by the accumulation of aggregated β-amyloid (Aβ), which is a 4-kDa peptide produced by the proteolytic cleavage of amyloid precursor protein (APP) by β- and γ-secretases. Oligomers of Aβ are the most toxic, whereas fibrillar Aβ is largely inert. Monomeric Aβ is found in normal patients and has an as-yet undefined function. The earliest consistent cytopathological change in Alzheimer disease is loss of synapses (1, 2). Finding a way to protect against the loss of synapses is a major therapeutic goal. Protein kinase C (PKC) activators have demonstrated neuroprotective activity in animal models of Alzheimer disease (3), depression (4), and stroke (5). Bryostatin, a potent PKC activator, also increases the rate of learning in rodents, rabbits, and invertebrates (4, 6, 7). This effect is accompanied by increases in levels of synaptic proteins spinophilin and synaptophysin and structural changes in synaptic morphology (8). PKC activators also can reduce the levels of Aβ and prolong the survival of Alzheimer disease transgenic mice (3). Evidence suggests that PKCε and ε are the most important PKC isoforms in eliciting these changes. Antisense inhibition of PKCε blocks secretion of sAPPα, whereas indirect activators of PKC, such as carbachol, increase sAPPα secretion (9). Experiments with specific PKC isoyme inhibitors also point to PKCe as the isozyme that most effectively suppresses Aβ production (10). Thus, isoform-specific PKC activators are highly desirable as potential anti-Alzheimer drugs. Specific activators are preferable to compounds such as bryostatin that show less specificity among conventional and novel forms of PKC because nonspecific activation of PKCε or β could produce undesirable side effects.

One compound known to activate PKCe specifically is DCP-LA (8-[(2-pentyl-cyclopropylmethyl)-cyclopropyl]-octanoic acid), a derivative of linoleic acid in which the double bonds are replaced by cyclopropane groups (11). Like the polyunsaturated fatty acids docosahexaenoic acid and arachidonic acid, DCP-LA binds to the phosphatidylserine-binding site and specifically activates PKCe (11). However, DCP-LA requires relatively high concentrations to produce its maximal effect. Therefore, we synthesized a variety of cyclopropanated fatty acid derivatives and compared their ability to activate PKCe. Two such compounds, DHA-CP6 and EPA-CP5, were found to be highly specific for the PKCe and were effective at 1000× and 100× lower concentrations than DCP-LA. DHA-CP6 also reduced the levels of Aβ by 60–70% in cells expressing APP with the Swedish mutation. This effect was not caused by activation of α-secretase, but may have been caused by increased Aβ degradation by endothelin-converting enzyme (ECE).

EXPERIMENTAL PROCEDURES

Materials—Culture media were obtained from K-D Medical (Columbia, MD) or Invitrogen. Aβ1–42 was purchased from AnaSpec (San Jose, CA). Polyunsaturated fatty acid methyl esters were obtained from Cayman Chemicals, Ann Arbor, MI. Other chemicals were obtained from Sigma-Aldrich.

Cell Culture—Human SH-SY5Y neuroblastoma cells (ATCC) were cultured in 45% F12K, 45% minimum Eagle’s medium, 10% fetal calf serum. Mouse N2a neuroblastoma cells were cultured in Dulbecco’s modified Eagle’s medium and 10% fetal calf serum without glutamine. Rat hippocampal neurons from 18-day-old embryonic Sprague-Dawley rat brains were plated on 24-well plates coated with poly-d-lysine (Sigma-Aldrich) in B-27 neurobasal medium containing 0.5 mm glutamine and 25 μm glutamate (Invitrogen) and cultured for 3 days in the

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2 The abbreviations used are: Aβ, β-amyloid peptide; APP, amyloid precursor protein; DAG, 1,2-diacyl-sn-glycerol; ECE, endothelin-converting enzyme; ELISA, enzyme-linked immunosorbent assay; MES, 4-morpholineethanesulfonic acid; PKC, protein kinase C; PMA, phorbol 12-myristate 13-acetate; TACE, tumor necrosis factor-α-converting enzyme; ADAM17, a disintegrin and metalloproteinase 17.
medium without glutamate. The neuronal cells were grown under 5% CO₂ for 14 days in an incubator maintained at 37 °C.

All experiments on cultured cells were carried out in triplicate unless otherwise stated. All data points are displayed as mean ± S.E. DCP-LA was used as its free acid in all experiments, whereas DHA-CP6, EPA-CP5, and AA-CP4 were used as their methyl esters.

**TACE—**TACE was measured (12) by incubating 5 µl of cell homogenate, 3 µl of buffer (50 mM Tris-HCl (pH 7.4) + 25 mM NaCl + 4% glycerol), and 1 µl of 100 µM TACE substrate IV (Abz-LAQAVRSSSR-Dpa) (Calbiochem) for 20 min at 37 °C in 1.5-ml polypropylene centrifuge tubes. The reaction was stopped by cooling to 4 °C. The samples were diluted to 1 ml, and the fluorescence was rapidly measured (ex = 320 nm, em = 420 nm) in a Spex Fluorolog 2 spectrofluorometer.

**PKC Assay—**After removal of culture medium, cells were scraped in 0.2 ml of homogenization buffer (20 mM Tris-HCl (pH 7.4), 50 mM NaF, 1 µg/ml leupeptin, and 0.1 mM phenylmethylsulfonyl fluoride) and immediately homogenized in the cell culture plate by sonication in a Marsonix microprobe sonicator (5 s, 10 W). Aliquots were transferred immediately after sonication to 0.5-ml centrifuge tubes and frozen at −80 °C. To measure PKC, 10 µl of cell homogenate or purified PKC isoforms were incubated for 15 min at 37 °C in the presence of 10 µM histones, 4.89 mM CaCl₂, 1.2 µg/µl phosphatidyl-L-serine, 0.18 µg/µl 1,2-dioctanoyl-sn-glycerol, 10 mM MgCl₂, 20 mM HEPES (pH 7.4), 0.8 mM EDTA, 4 mM EGTA, 8 µg/ml aprotinin, 8 µg/ml leupeptin, and 2 mM benzamidine. 0.5 µCi of [γ-32P]ATP was added, and [32P]phosphoprotein formation was measured by adsorption onto phosphocellulose as described previously (13). For measurements of activation by DCP-LA and similar compounds, PKC activity was measured in the absence of diacylglycerol (DAG) and phosphatidylserine, as described by Kanno et al. (11), and PKCδ and ε were measured in the absence of added EGTA and CaCl₂, as described by Kanno et al. (11). Low concentrations of Ca²⁺ are needed because high Ca²⁺ interacts with the PKC phosphatidylserine-binding site and prevents activation. For measurements of bryostatin activation, DAG was omitted unless otherwise stated. Freeze-thawing of the samples more than once was avoided because it was found to greatly reduce the PKC activity and the degree of activation.

**ECE Assay—**ECE was measured fluorometrically using the method of Johnson and Ahn (14). A sample of cell homogenate (20 µl) was incubated in 50 mM MES-KOH (pH 6.0), 0.01% C₁₂ E₁₀ (polyoxyethylene-10-lauryl ether), and 15 µM McaBk2 (7-methoxycoumarin-4-acetyl [Ala⁷-(2,4-dinitrophenyl)Lys⁹]-bradykinin trifluoroacetate salt) (Sigma–Aldrich). After 60 min at 37 °C, the reaction was quenched by adding trifluoroacetic acid to 0.5%. The sample was diluted to 1.4 ml with water, and the fluorescence was measured at ex = 334 nm, em = 398 nm.

**Other Assays—**Aβ1–42 human fluorometric ELISA kit (BioSource/Invitrogen). Aβ1 in culture medium was isolated by adsorption onto reversed phase columns (Waters Sep-Pak C₁₈ cartridges). The columns were conditioned with acetonitrile and equilibrated with water.

The sample was applied and washed with 2 ml of 5% acetonitrile and 2 ml of 25% acetonitrile containing 0.1% trifluoroacetic acid. The Aβ was eluted with 1 ml 50% acetonitrile + 0.1% trifluoroacetic acid. The acetonitrile was evaporated by heating under a stream of nitrogen, and the solution was lyophilized to dryness and redissolved in a minimum volume of water followed by ELISA measurements. Fluorescence was measured in a Bioteck Synergy HT microplate reader (ex = 460 nm, em = 560 nm). AlamarBlue and CyQuant NF (Invitrogen) were used according to the manufacturer’s instructions.

**Synthesis of Cyclopropanated Fatty Acids—**Methyl esters of polyunsaturated fatty acids were cyclopropanated using the modified Simmons-Smith reaction with chloroiodomethane and diethylzinc (15–17). All apparatus was baked at 60 °C for 1 h and flame-dried while passing dry nitrogen through the apparatus. A 100-ml three-neck round-bottom flask with a stirring bar and a temperature probe was surrounded by a dry ice mixture and filled with 1.25 g (4.24 mmol) of linoleic acid methyl ester or docosahexaenoic acid methyl ester in 25 ml of dichloromethane and bubbled with N₂. A 1 M solution of diethylzinc (51 ml, 54.94 mmol) in hexane was added anaerobically using a 24-inch-long 20-gauge needle, and the solution was cooled to −5 °C. Chloroiodomethane (CIC₂H₄I, 8.02 ml, 109.88 mmol) was added dropwise, 1 drop/s, with constant stirring. The rate of addition was decreased if necessary to maintain the reaction mixture below 2 °C. The reaction mixture became cloudy during the reaction, and an insoluble white zinc product was produced. The flask was sealed, and the mixture was allowed to react further for 1 h and then allowed to come to room temperature gradually over 2 h.

To prevent the formation of an explosive residue in the hood, diethylzinc was not evaporated off. The mixture was poured slowly into 100 ml of water under stirring to decompose any excess diethylzinc. Ethane was evolved. The mixture was centrifuged at 5000 rpm in glass centrifuge tubes, and the upper aqueous layer was discarded. The white precipitate was extracted with CH₂Cl₂ and combined with the organic phase. The organic phase was washed with water and centrifuged. The product was analyzed by Silica Gel G TLC using hexane–ethyl acetate (9 : 1) as the developing solvent. The starting material (16). Docosahexaenoic acid methyl ester was converted into DHA-CP6 in 90–95% yield. The product was a colorless oil with a single absorbance maximum at 202 nm in ethanol and no reaction with I₂. The IR spectrum showed cyclopropane ring absorption at 3070 and 1450 cm⁻¹. Under the same conditions, eicosapentaenoic acid methyl ester was converted to EPA-CP5, and arachidonic acid methyl ester was converted to AA-CP4. Linoleic acid methyl ester was converted to DCP-LA methyl ester, which was identical to a known sample.

**Hydrolysis of Methyl Ester—**The methyl ester (0.15 g) was dissolved in 1 ml of 1 N LiOH and 1 ml of dioxane. Dioxane and methanol were added until it became homogeneous, and the solution was stirred at 60 °C overnight to 3 days. The product was extracted in CH₂Cl₂ and centrifuged. The aqueous layer and
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FIGURE 1. Structures of DCP-LA, DHA-CP6, EPA-CP5, and AA-CP4.

white interface were reextracted with water and washed until the white layer no longer formed. The product was evaporated under N₂ and purified by chromatography on silica gel. The product, a colorless oil, eluted in 20% EtOAc in hexane. Its purity was checked by TLC in 10% EtOAc/hexane and by C18 reversed phase HPLC with UV detection at 205 nm, using 95% acetonitrile as the mobile phase.

RESULTS

The structures of the new PKC activators are shown in Fig. 1. To determine their PKC isozyme specificity, the new compounds were preincubated with purified isoforms of PKC for 5 min, and the PKC activity was measured radiometrically. DHA-CP6 and EPA-CP5, which are cyclopropanated derivatives of docosahexaenoic acid and eicosapentaenoic acid, preferentially activated PKCε to a similar extent. The maximal effect for DHA-CP6 and EPA-CP5 was observed at 10 nM and 100 nM, respectively (Fig. 2, A and B). AA-CP4, the cyclopropanated derivative of arachidonic acid, produced only modest activation (Fig. 2C), and cyclopropanated linolenyl and linoleyl alcohols, epoxyeicosatrienoic acid, and verencol acid methyl ester had little or no effect on PKC (Fig. 2E). Cyclopropanated verencol acid methyl ester inhibited PKCε at concentrations above 1 μM (Fig. 2E). DCP-LA also activated PKCε, as reported by Kanno et al. (11), but required 1000X higher concentrations than DHA-CP6 for a maximal effect (Fig. 2D). At higher concentrations, the compounds were weak inhibitors of PKCγ, δ, and βII.

Incubation of DHA-CP6, EPA-CP5, or AA-CP4 with cultured human neuroblastoma cells produced activation of PKC that remained elevated for up to 1 h (Fig. 3A). Sustained activation was even more pronounced in primary neurons, with PKC activation lasting up to 24 h (Fig. 3B). In contrast to conventional PKC activators such as phorbol ester, no PKC down-regulation to below control levels was observed. However, inhibition could be produced at extremely high concentrations (>100 μM) (Fig. 2). In contrast, PKC activators that bind to the PKC DAG-binding site, including bryostatin, gnidimacrin, and phorbol esters, produce a transient (5–10 min) activation of PKC activity, followed by a prolonged down-regulation (data not shown). Following application of PMA (phorbol 12-myristate 13-acetate) at its effective concentration of 10–100 nM, PKC is rapidly down-regulated and remains at 20–30% of control levels for up to 24 h, whereas down-regulation by bryostatin is milder and relatively short lasting (18–20). These differences have been attributed to differences in the DAG-binding C1 domain of PKC (19).

A growing body of evidence suggests that PKC down-regulation, particularly of PKCδ, may be responsible for the tumor-promoting effects of phorbol esters (18, 21). Down-regulation of PKC therefore complicates any proposed usage of DAG-binding PKC activators as therapeutic agents. Down-regulation has also been implicated as a factor in increasing Aβ production (22). Thus, the ability to produce sustained activation of PKC without inducing down-regulation is a highly desirable characteristic.

To measure the effects of PKCε activation on Aβ production, we used cells transfected with human APPsw/H1103, which produce large quantities of Aβ. Incubation of these cells for 24 h with PKC activators markedly reduced the levels of both intracellular (Fig. 4A) and secreted (Fig. 4B) Aβ. For bryostatin, which activates PKC by binding to the DAG-binding site, the inhibition was biphasic, with concentrations of 20 nM or higher producing no net effect. This may be explained by the ability of this class of PKC activators to down-regulate PKC when used at high concentrations. In contrast, DCP-LA and DHA-CP6, which bind to the PKC phosphatidylserine site, showed monotonically increasing inhibition at concentrations up to 10 to 100 μM with no evidence of biphasic activity at higher concentrations.

To determine whether the reduced levels of Aβ caused by PKC activators were due to inhibition of Aβ synthesis or activation of Aβ degradation, we applied DHA-CP6 and low concentrations (100 nM) of exogenous monomeric Aβ1–42 to cultured SH-SY5Y cells. This concentration of Aβ is too low to produce measurable toxicity or cell death. Because SH-SY5Y cells produce only trace amounts of Aβ, this experiment was an effective test of the ability of PKC activators to enhance Aβ degradation. By 24 h, most of the Aβ had been taken up by the cells, and the concentration of Aβ in the culture medium was undetectable. Addition of 0.01–10 μM DHA-CP6 to the cells reduced the cellular levels of Aβ by 45–63%, indicating that the PKCε activator increased the rate of degradation of exogenous Aβ (Fig. 5).

Previous researchers reported that PKC activators such as PMA produce large increases in TACE/ADAM17 activity (25, 26) which correlated with increased sAPPα and decreased Aβ, suggesting that TACE and BACE1 compete for availability of APP substrate and that PKC activators shift the competition in favor of TACE. However, many of these earlier studies were carried out in fibroblasts and other nonneuronal cell types, which may respond differently to PKC activators than neurons.

3 T. J. Nelson, unpublished results.
FIGURE 2. Activation of purified PKCε by PKC activators. Purified PKCα, βII, γ, δ, or ε (9 ng) was preincubated for 5 min at room temperature with DHA-CP6 (A), EPA-CP5 (B), AA-CP4 (C), or DCP-LA (D) followed by PKC activity measurement as described under “Experimental Procedures.” E, activation of purified PKCε by other cyclopropanated and epoxidized fatty acids, alcohols, and methyl esters. Samples were preincubated for 5 min at room temperature with purified PKCε (500 ng) followed by PKC activity measurement. Values are x ± S.E., two-tailed Student’s t test. *, p < 0.02; **, p < 0.001.
For example, Etcheberrigaray et al. (27) found that activation of PKC in human fibroblasts by 10–100 pM bryostatin increased the initial rate of ß-secretase activity by 16-fold and 132-fold, respectively. In contrast, in cultured human neuroblastoma cells (Fig. 6 A), PKC activators produced no detectable increase in TACE activity. A transient increase in TACE activity was observed in primary neurons after EPA-CP5 (Fig. 6 B). PMA also did not activate TACE in these cells (data not shown). This suggests that any reduction of ß-Amyloid by PKC activators is caused by some mechanism other than activation of TACE.

ß-Amyloid can be degraded in vivo by a number of enzymes, including insulin-degrading enzyme (insulysin), neprilysin, and ECE. Because PKCε overexpression has been reported to activate ECE (28), we examined the effect of PKC activators on ECE. Bryostatin, DCP-LA, and DHA-CP6 all produced a sustained increase in ECE activity (Fig. 7 A). A similar effect was observed in primary cultured neurons (Fig. 7 B). Because ECE does not possess a DAG-binding C1 domain or a PKC-like phosphatidylserine-binding C2 domain, this suggests that the activation was not due to direct activation of ECE, but must have resulted from indirect activation of ECE or some ECE-activating intermediate by PKC.

DHA-CP6, EPA-CP5, and AA-CP4 had no effect on the enzyme activity of purified ECE (Fig. 8).
and DCP-LA had no effect on cell survival or on proliferation as measured by alamarBlue and CyQuant staining (Fig. 9), indicating that the reduction in Aβ production did not result from cell proliferation or a change in cell survival.

**DISCUSSION**

DHA-CP6, EPA-CP5, and DCP-LA represent a new class of PKC activators that are relatively specific for the PKCe isoform. PKCe is expressed at very low levels in all normal tissues except for brain (29, 30). Wetsel et al. (31) reported that PKCe and PKCe are expressed predominantly in brain, with very weak staining in other tissues. The high abundance of PKCe in presynaptic nerve fibers suggests a role in neurite outgrowth and neurotransmitter release (32). Therefore, effects of specific PKCe activators would be largely restricted to brain and unlikely to produce unwanted peripheral side effects (18).

The concentration of DHA-CP6 (10 nM) required for activation of PKCe is comparable with that of PMA (10–100 nM) (33) but higher than that of bryostatin (0.2–1 nM). Further modifications of the structure of DHA-CP6, based on modeling studies of the PKC phosphatidylserine-binding site, could create derivatives with even higher affinity. Furthermore, because these new compounds are closely related to ω-3 and ω-6 polyunsaturated acids, they are likely to possess lower toxicity than many of the existing terpenoid secondary metabolite PKC activators. Thus, a requirement for higher concentrations might be less of a disadvantage.

There are several mechanisms by which activation of PKCe could affect Aβ production. One mechanism is by activating...
TACE/α-secretase and shifting the competition for APP away from the amyloidogenic β-secretase. This appears to be an important mechanism in nonneuronal cells such as fibroblasts; however, our data show that PKCe activators have much less effect on TACE activity in neuronal cell types than in fibroblasts. Specific inhibitors of PKCe also block PMA-induced suppression of Aβ levels (10), suggesting that PKCe is the principal isoform responsible for depleting Aβ. However, da Cruz et al. (22) found that long term application of phorbol ester increases the synthesis of PKCe. Thus, part of the effect could also be mediated by induction of PKCe by PMA. Decreased expression of PKCe also reduces cholinergic regulation of sAPPα levels (34), which is consistent with the theory that sAPPα and Aβ compete for a limited pool of APP. Thus, in cells in which APP is rate-limiting, direct or indirect activation of α-secretase by PKCe could also reduce Aβ levels. Another possible mechanism is by increasing the rate of degradation. A recent report (28) presented evidence that PKCe increases the activity of ECE, an enzyme that degrades Aβ. These authors reported that in APP/PKCε transgenic mice, which overexpress PKCe, levels of Aβ were lower than in control animals, but there was no change in other Aβ-degrading enzymes, such as insulin-degrading enzyme or neprilysin, and no change in APP metabolism. However, it was unclear whether this effect can be produced by activation of PKCe in nontransgenic, adult animals. Our results showing that Aβ degradation is enhanced in cells treated with specific PKCe activators suggest that this may be the case.

A third mechanism is by stimulation of PKC-coupled M1 and M3 muscarinic receptors, which is reported to increase nonamyloidogenic APP processing by α-secretase (TACE/ADAM17 and/or ADAM10) (35). Muscarinic agonists rescue 3× transgenic Alzheimer disease mice from cognitive deficits and reduce Aβ and tau pathologies (36), in part by activating the TACE/ADAM17 nonamyloidogenic pathway. Muscarinic receptor signaling is closely tied to PKC: muscarinic receptor mRNA is regulated by PKC (37), and neuronal differentiation produced by M1 muscarinic receptor activation is mediated by PKC (38).

Activation of PKC can also have other neurotrophic or synaptic effects that complement the effect of lowering Aβ. For example, PKC activators can reverse some of the neurological damage in an ischemia/hypoxia model for stroke (5). PKC activators are also reported to induce expression of tyrosine hydroxylase (39) and induce neuronal survival and neurite outgrowth (8, 40). These neurotrophic effects may be mediated by neurotrophins such as brain-derived neurotrophic factor (40, 41). Neurotrophin signaling and APP metabolism pathways intersect at many points. For example, p75 and TrkA nerve growth factor receptor signaling increase APP mRNA expression (42). Both APP and p75 are cleaved by α- and γ-secretase (43). Cleavage of the p75 extracellular domain enables fibroblast growth factor 2 to stimulate neurite outgrowth and branching (44).

An advantage of compounds such as DCP-LA which specifically activate PKCe is that, in contrast to phorbol esters and similar DAG analogs, they do not produce detectable down-regulation. The biphasic response of PKC to DAG-based activators means that a PKC activator may produce opposite effects at different time points, such as reducing Aβ levels at one time point and increasing them at another (e.g. Ref. 22). Careful dosing and monitoring of patients would be required to avoid effects opposite to those that are intended. Because of the relative inability of this new class of PKC activators to down-regulate PKC, this problem can be avoided.

FIGURE 8. Lack of activation of purified ECE by PKC activators. ECE activity was measured as described under “Experimental Procedures” in the presence of recombinant ECE (1784ZN, 20 ng; R&D Systems) and varying concentrations of DHA-CP6, EPA-CP5, or AA-CP4 for 60 min at 37 °C.

FIGURE 9. Lack of effect of DCP-LA and DHA-CP6 on cell survival and cell proliferation. Various concentrations of DCP-LA or DHA-CP6, or ethanol alone were added to SH-SY5Y cells growing on 6-well plates. Cell viability was measured by adding alamarBlue reagent (Invitrogen) to 10% of the culture volume. AlamarBlue is chemically reduced in viable cells to form a fluorescent product. The fluorescence was measured in a microplate reader at 1, 2, 3, and 4 h, and the slope of the curve of fluorescence intensity versus time was calculated. Cell proliferation was measured by adding 100 μl of CyQuant NF, which is a cell-permeable DNA-binding dye. After 60 min, the fluorescence, which is proportional to the number of cells, was measured in a microplate reader.
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