Increased Expression of Complement Decay-accelerating Factor during Activation of Human Neutrophils

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Abstract

Decay-accelerating factor (DAF) is a membrane protein that protects blood cells from damage by autologous complement. Using monoclonal antibodies in both direct-binding studies and flow cytometry, we found that resting neutrophils (polymorphonuclear leukocytes [PMN]) expressed 10^4 DAF molecules on their surface, and that surface DAF expression more than doubled when the cells were activated. Upregulation of surface DAF occurred within minutes, paralleled the upregulation of complement receptor types 1 and 3 (CR1 and CR3), and was not dependent on new protein synthesis. It was unaffected by EDTA but was inhibited by 10 μM trifluoperazine, suggesting involvement of intracellular Ca^{2+} and calmodulin or protein kinase C. Upon activation, the affected PMN lacking surface DAF from patients with paroxysmal nocturnal hemoglobinuria failed to increase DAF expression. In contrast, these cells increased CR1 and CR3 expression normally, suggesting that DAF deficiency in affected cells involves abnormal synthesis or packaging of DAF for intracellular storage. Translocation of DAF to the cell surface induced by chemoattractants may be important in allowing PMN to survive and function at inflammatory sites where there is rapid complement turnover.

Introduction

Decay-accelerating factor (DAF) of human erythrocytes (E) is a 70–75-kD surface protein which interacts with autologous C4b and C3b fragments when they become attached to the E membrane. By blocking uptake of the second component of complement (C), C2, and factor B, DAF inhibits assembly of the classical as well as the alternative pathway C3 and C5 convertases and thus interferes with the amplification steps of the complement cascade (1–3). DAF thereby prevents hemolytic activity from developing at sites where small numbers of C3b or C4b molecules become attached either directly to E, or indirectly to cells that are “bystanders” to the source of complement activation. Because DAF does not inhibit convertase formation on soluble immune complexes, neighboring cells, or foreign particles, it is believed that its physiologic function is to protect host E from damage by autologous complement (3). This hypothesis is strengthened by observations that the complement-sensitive E patients with paroxysmal nocturnal hemoglobinuria (PNH) are deficient in DAF (2, 4, 5), and that incorporation of purified DAF into the membranes of these cells ameliorates this abnormality (5).

Recent studies have demonstrated that DAF is also present on blood platelets, lymphocytes, monocytes, and neutrophils (polymorphonuclear leukocytes [PMN]) of normal individuals (6, 7). Comparison in a previous study of the content of DAF in cell extracts, as determined by immunoradiometric assay, with surface DAF expression, as determined by indirect immunofluorescence using monoclonal anti-DAF antibodies and flow cytometry, suggested that PMN contain more total DAF than is present on their surface membranes (6). These findings seemed analogous to results previously obtained for C3b and iC3b receptors (complement receptor types 1 and 3 [CR1 and CR3]), which have been shown to reside predominantly in intracellular pools in unstimulated circulating PMN (8–11). When the PMN are activated by chemoattractants, the intracellular CR1 and CR3 pools are rapidly translocated to the surface membrane by a process that does not require new protein synthesis (9–12). This translocation enhances the adherence and phagocytic activity of these cells as they move towards sites of infection (13).

Because increased amounts of “nascent” C4b and C3b molecules are likely to be present at inflammatory sites where complement activation is proceeding at a rapid rate, host PMN at these sites may be at increased risk for lysis by autologous complement. The present investigations were undertaken to determine whether the expression of DAF increases when PMN are activated. The results indicate that the amount of DAF expressed on the PMN surface rises at least twofold in response to complement activation products and other chemotactic stimuli. This increased DAF expression on PMN may be an important self-protective mechanism allowing these cells to survive and function while moving towards sites of complement activation in the inflammatory milieu. The results additionally indicate that in PNH, a disorder in which PMN are susceptible to uptake of increased amounts of autologous C3b (14) and because they are deficient in surface DAF (6, 15), affected cells are unable to augment DAF expression whereas they are able to increase CR1 and CR3 expression in a normal fashion. These results argue that synthesis or packaging of DAF is abnormal and that accelerated loss of DAF from the cell membrane is not the primary mechanism of DAF deficiency in affected PNH cells.

Methods

Patients. P.K. is a 56-yr-old woman with severe disease lasting 36 yr. She had undergone splenectomy in another city before the diagnosis of PNH was established by the acid lysis test. She has been transfusion dependent, requiring more than 2 U of washed red cells every 3 mo. She

Additional Details

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1. Abbreviations used in this paper: CR1 and CR3, complement receptor types 1 and 3; DAF, decay-accelerating factor; E, erythrocyte(s); FITC, fluorescein isothiocyanate; fMLP, N-formyl-methionyl-leucyl-phenylalanine; PNH, paroxysmal nocturnal hemoglobinuria; ZAS, zymosan-activated serum.

has had regional enteritis in addition to several complications of her PNH and is currently treated with iron dextran injections, daily folate, and 5-10 mg/d prednisone.

M.W. is a 30-year-old woman with anemia first recognized in 1982. She has had more mild disease, has not required routine transfusions, and is treated with daily folate and low doses of oral iron.

Complement lysis tests on blood samples from these patients were performed by Dr. Wendell Rosse, Duke University (16). P.K. was found to have 47% type III E (15-25-fold more sensitive than normal E), and M.W. was found to have 27% type II E (3-5-fold more sensitive than normal E).

Leukocyte isolation and incubation. PMN were isolated from freshly drawn heparinized blood of normal donors and patients as described previously (9). Briefly, the blood was layered over density gradients (17) of Percoll (Pharmacia Fine Chemicals, Piscataway, NJ) and centrifuged at 300 g at 14°C for 17 min. E were removed by two cycles of lysis using ice-cold 0.2% NaCl followed by addition of 1.6% NaCl. The leukocyte pellet was washed with Hank's balanced salts solution (HBSS) without Ca²⁺, Mg²⁺, or phenol red. The preparations generally contained >95% PMN of which >80% excluded trypan blue. The PMN fraction of patient P.K., with severe PNH, contained 10% E, which were larger and less dense than normal and some of which were nucleated. Leukocytes were obtained and purified as described previously (6). Fluorescein isothiocyanate (FITC)-conjugated F(ab')₂ fragments of goat anti-mouse IgG and anti-rat Ig were purchased from Cappel Laboratories, (Cochranville, PA). All antibodies and conjugates were centrifuged at 110,000 g for 15 min in an airfuge (Beckman Instruments, Inc., Fullerton, CA) before each use.

Immunofluorescent staining and flow cytometry. Immunofluorescent staining was carried out with all steps performed at 0-4°C (9). Briefly, after incubation, PMN were divided into aliquots of 10⁶ cells, washed with HBSS containing 0.1% bovine serum albumin, 0.05% NaN₃, and 10⁻³ M PMSF (HBSS+), and resuspended in 50 μl of the same solution. Monoclonal antibodies were added in excess as determined in preliminary titration experiments, and the mixtures were incubated on a rotator for 30 min. Cells were washed twice and resuspended in the same solution, excess FITC-F(ab')₂ anti-mouse IgG or anti-rat Ig conjugate was added, and incubation continued for another 30 min. The stained cells were washed three times, then resuspended in phosphate-buffered saline containing 0.05% NaN₃ and held on ice until examination in the flow cytometer.

The fluorescence of 10,000 cells from each sample was determined by flow cytometry using a Becton-Dickinson FACS Analyzer (Becton, Dickinson & Co., Sunnyvale, CA) standardized with 7.8-μm diameter fluorescent beads. The analyzer was operated with log-mode amplification, and the arithmetic mean linear equivalent fluorescence of each sample was computed automatically by the Consort 30 computer. Background fluorescence was determined using cells reacted with the FITC conjugate but no monoclonal antibody. This value was subtracted from all other determinations. Control experiments established that these background values did not differ from those obtained using isotype-matched, non-immune monoclonals in the first step (9).

Binding studies with 125I-anti-DAF. Purified IIH6 anti-DAF monoclonal was labeled with 125I using Iodogen (Pierce Chemical Co., Rockland, IL) to a specific activity of 6 × 10⁹ cpm/μg. Varying amounts of 125I-labeled antibody (0.01-0.20 μg) alone or with excess unlabeled antibody were diluted in a total volume of 0.1 ml of HBSS+ in 12 × 75-mm plastic culture tubes on ice. Aliquots of 0.1 ml of PMN which had been washed and suspended at 20 × 10⁶/ml after incubation were added, and the mixtures were vortexed briefly and incubated for 30 min at 4°C on a rotator. Duplicate 80-μl aliquots were removed from each tube, layered over 250 μl of synthetic silicone oil, d = 1.050 (General Electric Co., Waterford, NY) in 400-μl polypropylene centrifuge tubes and the cells were rapidly pelleted by centrifugation at 10,000 g for 2.5 min in a Beckman Instruments Microfuge B. 40 μl of the supernatant from each tube was counted to determine the amount of 125I-labeled antibody not bound. The remaining supernatant and most of the oil were aspirated and the tube was cut. The tip containing the cell pellet was counted to determine the bound radioactivity. Specific binding was determined by subtracting the radioactivity in the pellets of tubes containing 100-fold excess of unlabeled antibody from the corresponding tubes containing 125I-labeled antibody alone. Counts of duplicate aliquots from each incubation were averaged and corrected back to the original total volume, and the data was analyzed by the method of Scatchard (20) using an Apple IIe computer (Cupertino, CA).

DAF contents in NP-40 extracts of PMN were determined using an immunoradiometric assay as previously described (6). The absolute concentration of the DAF standard was established by amino acid analysis.

Results

Quantitation of surface DAF expression on isolated PMN and correlation with fluorescence measurements. In order to quantify the number of DAF molecules on the surface of PMN, increasing amounts of 125I-anti-DAF monoclonal antibody IIH6 were incubated with the cells and both total and specific binding was determined. Results of a representative experiment are shown in Fig. 1 A and the corresponding Scatchard analysis is shown in Fig. 1 B. In this study, which was performed on freshly isolated cells held in the resting state at 0°C, the apparent dissociation constant was 1.6 × 10⁻⁹ M and the estimated number of sites

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Quantitation of surface DAF on PMN by binding of 125I-anti-DAF. (A) The results shown are the average of duplicate aliquots from the incubation mixtures at each input of labeled antibody. Non-specific binding was determined in the presence of a 100-fold excess of unlabeled antibody. Specific binding = total − non-specific binding. All counts were corrected by subtracting background. (B) Scatchard analysis of data from A. By linear regression, Kd = 1.64 × 10⁻⁹ M and extrapolated binding at saturation = 8,540 anti-DAF molecules per cell with r = 0.99.
at saturation was \(8.5 \times 10^8\)/cell. The mean of the dissociation constants calculated from eight similar studies was \(5.8 \times 10^{-9}\) M.

To permit conversion of relative fluorescence intensity data obtained using flow cytometry to numbers of molecules per cell, immunofluorescent and direct-binding determinations were carried out in parallel on aliquots of cells from the same incubation mixtures. Monoclonal anti-DAF IA10 and FITC-conjugated F(ab)\(_2\) anti-mouse IgG were used for the fluorescent analysis and \(^{125}\)I-IIH6 for the direct-binding determinations. As shown in Table I, the mean (±SEM) level of DAF expression in resting cells as determined by the direct-binding method was 9,740±1,070 molecules per cell, and in activated cells was 22,700±720 molecules of DAF per cell. The comparative ratios of surface DAF on activated to resting cells obtained with relative fluorescence measurements, 2.58, and direct binding determinations, 2.33, were in close agreement. When subjected to linear regression analysis, the correlation coefficient (\(r\)) was 0.97.

**Surface DAF expression on isolated PMN held at 0°C and effect of temperature.** Previous studies of isolated PMN held at 0°C have shown that resting PMN express \(\approx 6,000\) CR1 and 17,000 CR3 molecules/cell (8, 9). Studies of isolated PMN at 0°C using anti-DAF monoclonals are shown in the left group of bars in Fig. 2 in relation to control studies using anti-CR1 and anti-CR3 monoclonals. As can be seen, higher fluorescence on resting cells at 0°C was obtained with anti-DAF than with the antibodies to either of the complement receptors. Because CR1 and DAF were stained with the same FITC-conjugate, the relative fluorescent intensities reflect the relative numbers of each protein, assuming one antibody molecule binds to each protein molecule (6, 9, 12). The value obtained for DAF, 3.53±0.38, (mean±SEM, \(n = 7\)) was 2.2 times higher than that for CR1, 1.59±0.51 in this series of determinations.

The effect of temperature on DAF expression on the isolated PMN was evaluated next. As shown in Fig. 2, surface DAF levels increased 1.47-fold when the cells were warmed from 0°C to 37°C. This increase resembled the increases in CR1 and CR3 expression which have been observed when buffy coat cells or PMN isolated by centrifugation are warmed to 37°C in the absence of chemical stimuli (8–10). In the determinations of CR1 and CR3 expression performed here in parallel to the analyses of DAF, the mean levels of the two receptors increased 2.75- and 4.56-fold, respectively.

**Increased surface DAF expression induced by chemoattractants.** DAF levels on PMN after stimulation with fMLP or ZAS were next measured as shown in the right half of Fig. 2. Compared with cells held at 37°C in medium alone, cells treated with \(10^{-8}\) M fMLP showed a further 2.13±0.48-fold increase in DAF expression. Binding studies with \(^{125}\)I-anti-DAF monoclonal IIH6 demonstrated that this level of fluorescence corresponded to 22,700±720 DAF molecules per cell (mean±SEM, \(n = 3\)). Cells treated with ZAS, a source of C5a des-Arg, showed a 2.08±0.15-fold increase in DAF fluorescence. Quantitation of total DAF in extracts of similar cell preparations by immunoradiometric assay revealed 18,000±1,000 (mean±SEM, \(n = 8\)) molecules per cell, with an additional 1,800–3,000 molecules per cell remaining insoluble. In the same experiments, fMLP induced 3.82- and 3.41-fold increases in CR1 and CR3 expression whereas ZAS increased surface CR1 and CR3 3.47- and 3.46-fold, respectively.

The time course and dose response for the effects of fMLP on DAF expression are shown in Fig. 3. DAF expression increased to 80% of maximum within 10 min after addition of \(10^{-8}\) M fMLP. Cells examined 60 min after addition of the peptide continued to show near-maximal DAF expression. \(10^{-10}\) M fMLP had no discernable effect whereas \(10^{-8}\) M gave about half of the maximal response, which was observed at \(10^{-8}\) M. Higher concentrations caused \(\approx 10\%\) less DAF expression. The dose-response curves for CR1 and CR3 generated in the same experiments were parallel to that shown for DAF.

**Effect of protein synthesis inhibitors and possible role of Ca\(^{2+}\) in augmenting DAF expression.** The rapidity with which the expression of DAF increased in response to the above stimuli, together with the previous demonstration of a large excess of

![Figure 2. Comparison of DAF, CR1, and CR3 expression on resting and stimulated isolated PMN. Cells were incubated under the designated conditions for 1 h before washing and immunofluorescent staining. Background fluorescence was subtracted from all values as described in the Methods. Results shown are the means±SEM of the fluorescence of 10,000 cells determined in each of seven experiments. The fMLP concentration was \(10^{-8}\) M and ZAS was used at 1:100 final dilution. Temperatures are given in °C.]

| Table I. Quantitation of DAF Expression on Resting and Activated Isolated PMN |
|---------------------------------|-----------------|-----------|
| Mean fluorescent                | \(^{125}\)I-anti-DAF | n         |
| fluorescence                    | sites/cell       |           |
| Resting: 0°C in                 |                 |           |
| HBSS/gel                       | 5.99±0.34       | 9,740±1,074 | 4       |
| Stimulated: fMLP                | 15.44±0.90      | 22,700±719  | 3       |
| 10\(^{-8}\) M                   |                 |           |
| Ratio of means:                 |                 |           |
| Stimulated/Resting              | 2.58            | 2.33      |

Aliquots of each cell suspension were split: 2 \(\times\) \(10^6\) cells were used for fluorescence analysis and 24 \(\times\) \(10^6\) were used for direct binding study. Results shown are mean±SEM for the number of experiments indicated. Fluorescence was corrected for background and autofluorescence using FITC conjugate but without monoclonal anti-DAF. Scatchard analysis of \(^{125}\)I-binding data was performed as in Fig. 1. Linear regression analysis of mean fluorescence vs. \(^{125}\)I binding had \(r = 0.97\).
Increased Decay-accelerating Factor Expression during Neutrophil Activation

DAF antigen in PMN extracts not detectable on the PMN surface suggested that augmented DAF expression might involve translocation of DAF from an intracellular pool rather than new DAF synthesis. We therefore determined the effects of protein synthesis inhibitors. The increase in DAF expression induced by fMLP, 197±6% (n = 5) was not altered by the addition of 10 µg/ml of cycloheximide, 203±14% (n = 3), or puromycin, 202±13% (n = 4). Similar results were obtained with ZAS. This is consistent with the hypothesis that new protein synthesis is not required.

The effects of agents which modulate extra- and intracellular Ca²⁺ levels were investigated next. The results shown in Fig. 4 indicate that adding neither 1.2 mM CaCl₂ nor 5 mM EDTA to media initially free of divalent cations had any effect on the augmentation of DAF expression by fMLP. In contrast, 10 µM trifluoperazine caused significant inhibition (P < 0.01), suggesting a role for intracellular Ca²⁺ and calmodulin and/or protein kinase C. To further evaluate this possibility, ionomycin, an agent which bypasses the physiologic signal-transducing mechanisms, was used to increase the intracellular Ca²⁺ concentration directly. As shown by the solid bars in Fig. 4, an increase in DAF expression resulted that was comparable to the increase induced by fMLP. In contrast to the fMLP effect, however, the effect of ionomycin was dependent on extracellular Ca²⁺ and was markedly inhibited by EDTA. These results parallel those previously reported for CR1 (21) and suggest that increased expression of DAF may similarly depend on the release of Ca²⁺ from intracellular stores rather than on the influx of extracellular Ca²⁺ which is also initiated by fMLP.

**DAF expression on PMN from patients with PNH.** Recent studies have shown that DAF is deficient in affected PMN of PNH patients (6, 15). In view of the above observations that a major portion of DAF in circulating PMN resides inside the cell rather than on the membrane, we next investigated whether PNH PMN could increase their DAF expression in response to activating stimuli. FACS Analyzer histograms of the immunofluorescence of PMN of a normal donor and two PNH patients are shown in Fig. 5. As seen in Fig. 5 A, the PMN from the normal donor appeared as a single population with respect to DAF, CR1, and CR3 staining, and the expression of all three surface proteins increased in response to stimulation with fMLP. Fig. 5 B shows parallel studies on PMN from a patient (M.W.) with type II E. Staining for DAF revealed two distinct populations of PMN, one with no fluorescence above background, and another which appeared to express a relatively normal amount of DAF in the resting state. When these cells were stimulated with fMLP, the DAF-negative population did not change while the mean fluorescence of the DAF-positive population increased from 13.9 to 20.1. When corrected by subtracting the background fluorescence, this change represented a 1.6-fold increase, approximating that shown with PMN from normal individuals (see above). In contrast to the findings for DAF, this individual’s PMN appeared as one homogeneous population with regard to CR1 and CR3 expression, and all cells increased these markers normally in response to stimulation. The results with PMN from a patient (P.K.) with type III E are shown in Fig. 5 C. In distinction to patient M.W., with type II E described above, the resting PMN showed a single population with DAF fluorescence indistinguishable from background. There was little or no change in DAF expression when these cells were stimulated. As was the case with PMN from patient M.W., these cells appeared as a single population with regard to CR1 and CR3 expression, and the expression of both receptors increased normally after stimulation. The latter histograms show a small percentage of contaminating large erythrocytes which were gated out and were revealed as negative when the CR1 and CR3 increased. Taken together, these results suggest that affected PMN in PNH not only lack surface DAF but are unable to increase DAF expression in response to activating stimuli.

**Discussion**

In this report, we present evidence that PMN contain intracellular DAF and that some of this DAF is rapidly brought to the PMN surface when the cells are activated. We found that 10⁴ DAF molecules are present on the surface of isolated PMN held at 0°C, and that there is a small spontaneous increase in DAF
expression when isolated cells are subsequently placed at 37°C in the absence of stimuli. An additional twofold increase in DAF expression is induced by treatment with fMLP or with ZAS, a source of C5a des Arg. Maximal DAF expression occurs in minutes and is not inhibited by cycloheximide or puromycin. Thus it does not seem to involve protein synthesis. The maximum amount of DAF expressed on the cell surface in these studies, about 23,000 molecules per cell, corresponds to the total content of DAF in cell extracts. It thus seems likely that most or all of the intracellular DAF is available for translocation to the PMN surface in response to stimulation by optimal doses of fMLP or other activating agents.

It has been shown previously that, when human PMN are stimulated by chemotaxants or other activators, there is a
rapid increase in the expression of CR1 and CR3, the receptors for the major opsonically active components of the complement system (8–13). The number of receptors for formyl-peptide chemoattractants also increases under these conditions (22). In addition to the appearance of these receptors on the membrane, there is an increase in the association with the membrane of protein kinase C activity and of cytochrome b245, a component of the microbicidal oxidase system (23, 24). These alterations in membrane protein expression are specific, however, inasmuch as there is no concomitant change in the number of Fcγ receptors or in β-2-microglobulin expression (9). Thus, simple unfolding or ruffling of redundant membrane cannot account for these phenomena which are believed to be of critical importance in optimizing the function of the activated cell.

The intracellular pools of CR1 and CR3 are distinct from each other (11). Optimal CR3 expression is inhibited by EDTA and Mg-EGTA, whereas optimal CR1 expression requires intracellular Ca2+ but is not inhibited by these chelators (21), indicating that some of the reactions responsible for translocation of the two distinct receptor proteins to the surface are also apparently distinct. The characteristics of upregulation of DAF expression in response to chemoattractants are thus similar to those for CR1. The data suggest that induction of this translocation may involve intracellular Ca2+, in that this phenomenon is blocked by 10 μM trifluoperazine, which inhibits calmodulin and/or protein kinase C-dependent processes (25, 26), but not by EDTA. Further evidence suggesting a role of Ca2+ is provided by the increase in DAF expression induced by ionomycin with Ca2+. There appears to be more DAF than CR1 on the surface of isolated PMN maintained at 0°C as judged both by relative fluorescence and by direct-binding studies of 125I-anti-DAF monoclonal antibodies. Recent studies have shown that E DAF differs from CR1 in that it is anchored to the membrane by a C-terminal glycolipid structure that is added to the DAF molecule during posttranslational processing (27). Determination of the relationship of the intracellular pool(s) of DAF to those of CR1 and CR3 and elucidation of the precise mechanism by which DAF is translocated to the surface membrane await further study.

Although the underlying defect in PNH is not yet clarified, the deficiency of DAF in E plays a major role in pathophysiology (1–7). Reconstitution of PNH E with DAF restores their ability to circumvent autologous C3b uptake and subsequent lysis (5). Immunohistochemical analyses have demonstrated that DAF is deficient in PMN of all PNH patients studied to date (4, 6). In the two PNH patients described in this study, this was also the case, and the percentage of DAF-negative circulating PMN was much greater in the patient with type III E. It is possible that circulating PMN of PNH patients could lose surface DAF while retaining their intracellular pool or that affected PMN could generate new DAF upon activation. The results clearly indicate that DAF expression on initially negative cells does not increase when they are stimulated. In contrast, both CR1 and CR3 expression on the DAF-deficient cells appear to increase normally. Interestingly, in the patient whose PMN showed two distinct populations when stained for DAF, the same cells appeared homogeneous when stained for CR1 and CR3, and the population of initially DAF-positive PMN further increased DAF expression normally after stimulation. Thus, circulating PMN that are deficient in surface DAF also appear to be deficient in intracellular DAF, suggesting that the deficiency occurs at the time of synthesis or packaging of DAF for intracellular storage. Accelerated loss of DAF from the membrane is therefore unlikely to account for the deficiency of DAF in affected cells.

Because PMN must migrate to and function in inflammatory sites where complement consumption is proceeding at rapid rates, it seems advantageous that the expression of DAF on their membranes should increase in response to chemoattractants. The major function of DAF is believed to be protection of the cell against damage by autologous complement. Such damage could certainly occur if increased amounts of C4b and C3b, generated by convertases on other cells and/or in the fluid phase, became deposited on the infiltrating host PMN, and served as sites for formation of additional convertase activity. This amplification is specifically inhibited by DAF. The increased expression of DAF under these conditions may thus be important in allowing the PMN to survive and function at sites of infection and inflammation.

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