Components of Conidial and Hyphal Walls of
Penicillium chrysogenum

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(Accepted for publication 30 June 1969)

SUMMARY

Cell walls were prepared mechanically from both hyphae and conidia of
Penicillium chrysogenum. The major carbohydrate components of cell wall
hydrolysates were glucose, galactose and glucosamine, with lesser amounts of
mannose and rhamnose. The conidia were richer in galactose and the hyphae
contained more glucosamine. The distribution of the monomers in acid-
soluble and alkali-soluble fractions was different in the two kinds of cell wall;
conidial walls contained higher concentrations and a greater variety of
amino acids than hyphal walls.

INTRODUCTION

In contrast to the intensive investigations of the polymers constituting the cell walls
of bacteria, the study of the structural macromolecules of the cell walls of filamentous
fungi has been very limited. Many of these studies, dating from the last century, were
restricted to histological observations (van Wisselingh, 1898; Melzer, 1924). More
recent reports have been concerned with direct chemical analyses. In most cases
(Hamilton & Knight, 1962; Crook & Johnston, 1962; Manocha & Colvin, 1967;
Applegarth, 1967), however, analyses have been restricted to only the hyphal stage in
the developmental cycle, although the acetyl content of hyphal and spore walls of
Asperigillus phoenicis has been compared (Bloomfield & Alexander, 1967).

Penicillium chrysogenum, a filamentous fungus, possesses at least two morphologi-
cally distinct stages representing a seemingly primitive morphogenetic system. The
conidia are oblate spheroids, green in colour, and possessing rigid cell walls, whereas
the hyphae which develop from them are filamentous, flexible and white. These
morphological differences suggest differences in chemistry. Mahadevan & Tatum
(1965) have detected a number of quantitative differences in the cell walls of a variety
of mutants of Neurospora crassa. They concluded that the morphology is a direct
consequence of the cell wall composition and that changes from wild-type growth to
other forms may be caused by alterations in the relative concentrations of the struc-
tural polymers of the cell wall. It seems reasonable to expect that functionally com-
parable structures which differ morphologically at different stages in the life cycle of an
organism would also differ chemically. Because of the paucity of comparative data and
to uncover basic molecular changes in the conversion of conidium to hyphal cell,
studies were undertaken to determine whether the cell walls of the two stages are, in
fact, biochemically distinct.

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METHODS

Organism and growth conditions. Penicillium chrysogenum, strain NRRL 1951-B 25, was maintained on Sabouraud glucose (SG) slopes: 4% glucose, 1% 'neopeptone', 1.8% agar. After sporulation, the cultures were refrigerated.

Roux culture bottles containing SG agar were inoculated with conidia from sporulated slopes and incubated at ambient temperature (about 25°C) until sporulation occurred, which was usually 4 days. Sterile water was added, and the conidia removed by scraping the surface with a sterile aluminium spatula. The conidial suspension was filtered through sterile Pyrex glass wool so that the spore harvest would be free from mycelial fragments, as verified by microscopic examination. After at least five cycles of washing and centrifugation, the conidia were suspended and maintained in cold M/15 potassium phosphate buffer (pH 7.4).

Hyphae were grown by inoculating a spore suspension into 6 l. of SG broth in a Microferm Laboratory Fermentor (New Brunswick Scientific Co., Inc., New Brunswick, New Jersey, U.S.A.). Incubation continued for 48 hr at ambient temperature under constant aeration and agitation. The hyphae were collected on filter paper and washed four times with cold distilled water.

Preparation of spore walls. Washed spores (10 g. wet weight) were suspended in 10 ml. cold phosphate buffer (pH 7.4) to which 50 g. of glass beads (0.5 mm. diameter) were added. Spores were disrupted in a Bronwill Mechanical Cell Homogenizer maintained at 5°C. When disruption was essentially complete (as determined by the release of protein into the supernatant fluid and verified microscopically), homogenization was discontinued. The mixture was centrifuged at 500 g to separate the spore walls from the glass beads. The crude wall fraction was collected by centrifugation at 4000 g. The pellet was suspended in 100 volumes of distilled water and agitated on a wrist action shaker overnight at 4°C. The suspension was centrifuged and washed successively with distilled water and phosphate buffer (pH 7.4) until no extinction was detectable in the supernatant at 260 mp and 280 mp, indicating the absence of cytoplasmic contaminants. After collection, the pellet was extracted for 1 hr in each of the following solvents: 95% (V/V) ethanol in water at 5°C, three times with boiling ethanol + ether (1:3, V/V), and finally boiling ether. After extraction the walls were lyophilized.

Preparation of hyphal walls. Hyphae were disrupted in a Sorvall Omnimixer (Ivan Sorvall, Inc., Norwalk, Connecticut, U.S.A.) with glass beads. Purification procedures were the same as described for the preparation of spore walls.

Isolation of carbohydrates. The procedure described by Mahadevan & Tatum (1965) was adopted: 49 mg. samples of wall material were treated with 2 N-NaOH for 16 hr at room temperature, centrifuged, and the supernatant liquid treated with 2 vol. 95% (V/V) ethanol in water. The resulting precipitate was suspended in water, dialysed against water, and lyophilized; this fraction was designated F1. The residue remaining after the treatment with NaOH was resuspended in N-sulphuric acid, incubated at 90°C for 16 hr and centrifuged. The pellet was washed and the washings were added to the supernatant fluid. The supernatant fluid was adjusted to pH 7 with Ba(OH)2, and the resulting precipitate was removed by centrifugation and discarded; the material in the supernatant fluid was designated F2. The residue remaining from the sulphuric acid treatment was resuspended in 2N-NaOH for 30 min. at room temperature and
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centrifuged and 2 vol. ethanol added to the supernatant liquid. The precipitate was centrifuged down, suspended in water, dialysed against water, and lyophilized; this fraction was designated F3. The residue from the second NaOH treatment was suspended in water, dialysed, lyophilized, and designated F4. Before electrophoresis, the fractions were hydrolysed according to the procedure of Mahadevan & Tatum (1965).

**Extraction of amino acids.** Samples (50 mg.) of either conidial or hyphal walls were heated in 6N-HCl in evacuated tubes at 105° for 18 hr. The hydrolysates were filtered through 'Millipore' filters. The filtrates were taken to dryness by heating in vacuo in a hot water bath. Each hydrolysate was dissolved in 1 ml. 10% (v/v) isopropanol in water and stored at 0°.

**Separation of components.** Separation of amino acids was accomplished by electrophoresis with the high voltage electrophorator, Model D (Gilson Medical Electronics, Middletown, Wisconsin, U.S.A.). Hydrolysate samples or standards were applied at 5 cm. intervals parallel to the short side along an origin line drawn 12·7 cm. from the end of a 46·4 x 57·2 cm. sheet of Whatman 3 MM chromatography paper. The paper was then sprayed with a mixture of formic acid + acetic acid + water (6 + 24 + 170 by vol.) at pH 2 until wet and transferred to the electrophoresis tank with the origin line proximal to the anode. The samples were run in the same buffer at 30 V/cm., current 80 to 85 mA, for 90 min. at 10 to 12°. After electrophoresis, the paper was dried at 90° for 20 min. Neutral sugars were run in a boric acid buffer adjusted with NaOH to pH 9·4 at 240 V/cm., current 4 to 5 mA, for 150 min. at 10 to 15°. After removal from the electrophoresis chamber, the paper was dried as described above. Because of trailing, the resolution obtained with amino sugars upon direct electrophoresis was insufficient for identification or quantitation, so the samples were oxidized with ninhydrin before electrophoresis; this converted any hexosamines present to their corresponding pentoses (Stoffyn & Jeanloz, 1954). The oxidized samples were run as described for neutral sugars.

Amino acids were made visible by dipping the electrophoretograms in 1·5% ninhydrin in 95% (v/v) ethanol in water, drying at room temperature and counterstaining with a cupric nitrate solution (Mabry & Todd, 1963). Sugars were made visible with the diphenylamine-p-anisidine reagent (Mabry, Gryboski & Karam, 1965).

In situations where components could not be separated by electrophoresis, the electrophoretograms were cut into strips parallel to the direction of migration. The strips were sewn on to Whatman No. 1 paper and chromatographed in the second dimension with 2-butanone + propionic acid + water (75 + 25 + 30, by vol.). By this procedure, amino acids which had similar electrophoretic mobilities were separated and amino acids whose mobility properties left them within the glucosamine streak were resolved.

**Identification and estimation of components.** The amino acids and sugars were identified by three properties: (1) electrophoretic mobility compared with standard samples run simultaneously; (2) characteristic tints after colour development; (3) rates of colour development. Electrophoretic mobility values were also compared with published values (Mabry & Todd, 1963; Himes & Metcalfe, 1959).

Quantitative estimates were made by cutting the developed sheets into strips 3·2 cm. wide and scanning them in a Model RB Analytrol densitometer-integrator (Spinco Division, Beckman Instruments, Inc., Palo Alto, California, U.S.A.). For the amino
acids, 500 mμ interference filters were used, and for the sugars, 450 mμ interference filters. Proportionality between integration units and concentration was obtained in the range 0 to 400 μg. for the sugars and 0 to 100 μg. for the amino acids.

**Nitrogen analysis.** Nitrogen was determined as ammonia by the Kjeldahl-Nessler method described by Wilson & Knight (1952).

**RESULTS**

The distribution of monosaccharides in hydrolysates of the series of extracts from conidial and hyphal cell walls is shown in Table 1. The first alkali-soluble fraction (F 1) of conidial walls contained glucose, galactose and mannose. Hydrolysis of hyphal F 1 also gave glucose and galactose, but no mannose. Quantitatively, the two fractions differed markedly in the relative amounts of glucose and galactose. The glucose:galactose ratio of conidial F 1 was 0.32, whereas in hyphal F 1, the ratio was 7.5. The total weight of the two fractions were similar, conidial F 1 being slightly greater.

<table>
<thead>
<tr>
<th></th>
<th>Conidial walls</th>
<th></th>
<th>Hyphal walls</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>F1/mg./100 mg.</td>
<td>F2/mg./100 mg.</td>
<td>F3/mg./100 mg.</td>
<td>F4/mg./100 mg.</td>
</tr>
<tr>
<td>Glucose</td>
<td>6.12</td>
<td>5.92</td>
<td>14.30</td>
<td>26.34</td>
</tr>
<tr>
<td>Galactose</td>
<td>18.80</td>
<td>0.61</td>
<td>19.41</td>
<td>7.54</td>
</tr>
<tr>
<td>Mannose</td>
<td>2.25</td>
<td>0.33</td>
<td>2.58</td>
<td>1.00</td>
</tr>
<tr>
<td>Rhamnose</td>
<td>0.31</td>
<td></td>
<td>0.31</td>
<td>0.13</td>
</tr>
<tr>
<td>Glucosamine</td>
<td>4.08</td>
<td></td>
<td>7.35</td>
<td>11.43</td>
</tr>
<tr>
<td>Residue</td>
<td>6.95</td>
<td></td>
<td>11.02</td>
<td>17.97</td>
</tr>
<tr>
<td>Total</td>
<td>34.12</td>
<td>11.25</td>
<td>14.30</td>
<td>18.37</td>
</tr>
</tbody>
</table>

Both F 1 fractions were non-dialysable and could be precipitated from NaOH with ethanol or ammonium sulphate. However, the conidial F 1 was dark green, which was not the case with hyphal F 1. Despite the intense colour, no extinction bands were detectable in a scan of the material in the visible spectrum. The extinction curve obtained was suggestive of light scattering of sufficient intensity to mask any specific absorption bands. On standing at room temperature for extended periods of time, the colour passed through a dull brown colour to a mustard yellow. This colour change was accelerated by reducing agents, e.g. sodium dithionite, and reversed by hydrogen peroxide. The hyphal F 1, in contrast, did not exhibit these oxidation-reduction characteristics. The nature and function of the green pigment is unknown.

The F 2 fractions of conidial and hyphal walls contained all five detectable monosaccharides. However, they differed quantitatively, each monosaccharide being present in higher concentration in the hyphal walls. Consequently, hyphal F 2 constituted a significantly greater fraction (30%) than conidial F 2 (11%).

The two fractions F 3 and F 4 appeared to be the only homopolymeric fractions. On hydrolysis, F 3 gave only glucose, and F 4 only glucosamine. There were about the same amounts of glucosamine in the F 4 fractions of either wall. The conidial F 3 contained twice as much glucose as in hyphal F 3.

Comparison of the total amounts of each monosaccharide indicates that only galactose and glucosamine differed markedly in the two kinds of wall. The galactose...
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The content of the conidial walls was 2.7 times that found in the hyphal walls. The hyphal walls, on the other hand, were richer in glucosamine, containing 1.6 times that found in conidial walls. Glucose was present at slightly greater concentration in the hyphal walls. Although the rhamnose content of hyphal walls was greater than in conidial walls, the concentrations in both kinds of wall was sufficiently low so that rhamnose may be described as a minor component.

Other differences in the composition of the cell walls became apparent from the distribution of the monosaccharides in the several fractions. All of the mannose of hyphal walls was in the F2 fraction, whereas virtually all appeared in fraction F1 of conidial walls. Although glucose was found in fractions F1, F2 and F3 of both types of walls, it was concentrated in F3 in conidial walls, whereas it was present in largest amounts in F1 and F2 of hyphal walls. Essentially all of the galactose of conidial walls was recovered in F1, but in hyphal walls it was found in F1 and in F2, with 70% of the total galactose in F2. Glucosamine was found at about the same concentration in F4 of both hyphal and conidial walls. However, hyphal F2 contained significantly more glucosamine than did conidial F2.

Included in Table 1 are mole ratios for monosaccharide components of *Penicillium chrysogenum* hyphal walls reported by Hamilton & Knight (1962). The present results appear to be in good agreement with those data for galactose, mannose and rhamnose, but are high with respect to glucose and glucosamine. In the paper by Hamilton & Knight (1962), xylose was reported as a component of the hyphal walls; this pentose has also been reported as a trace component in *P. digitatum* and *P. italicum* (Grisaro, Sharon, & Barkai-Golan, 1968). Although the presence of xylose had been expected, we found no xylose in the hydrolysates of any fraction. Neither Crook & Johnston (1962) nor Applegarth (1967) detected xylose in hydrolysates of the closely related *P. notatum*. Since the organism used in the present work is the strain Hamilton & Knight (1962) used, it seems likely that the absence of xylose had resulted from differences in growth conditions.

### Table 2. Amino acid components of conidial and hyphal walls of *Penicillium chrysogenum*

<table>
<thead>
<tr>
<th>Amino acid</th>
<th>Conidial walls (mg./100 mg. wall)</th>
<th>Mole ratio</th>
<th>Hyphal walls (mg./100 mg. wall)</th>
<th>Mole ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aspartic acid</td>
<td>3.92</td>
<td>1.73</td>
<td>0.61</td>
<td>3.40</td>
</tr>
<tr>
<td>Glutamic acid</td>
<td>2.78</td>
<td>1.10</td>
<td>0.29</td>
<td>1.46</td>
</tr>
<tr>
<td>Threonine</td>
<td>0.25</td>
<td>0.12</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Leucine</td>
<td>0.83</td>
<td>0.38</td>
<td>0.22</td>
<td>1.17</td>
</tr>
<tr>
<td>Isoleucine</td>
<td>0.83</td>
<td>0.38</td>
<td>0.20</td>
<td>1.13</td>
</tr>
<tr>
<td>Valine</td>
<td>1.22</td>
<td>0.61</td>
<td>0.20</td>
<td>1.27</td>
</tr>
<tr>
<td>Serine</td>
<td>0.67</td>
<td>0.38</td>
<td>0.13</td>
<td>0.92</td>
</tr>
<tr>
<td>Alanine</td>
<td>1.51</td>
<td>1.00</td>
<td>0.12</td>
<td>1.00</td>
</tr>
<tr>
<td>Glycine</td>
<td>1.47</td>
<td>1.15</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Arginine</td>
<td>1.88</td>
<td>0.64</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Lysine</td>
<td>1.58</td>
<td>0.64</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Total</td>
<td>16.94</td>
<td>—</td>
<td>1.77</td>
<td>—</td>
</tr>
</tbody>
</table>

The amino acid composition of the two types of wall differed not only in total amount and relative proportions, but qualitatively as well (Table 2). No tyrosine, phenylalanine, tryptophan, cysteine, methionine or histidine was detected in the conidial or hyphal walls. Furthermore, hyphal walls did not contain any glycine,
threonine, lysine or arginine, all of which were found in conidial walls. The absence of these amino acids raised the question of their possible destruction during hydrolysis. There have been reports of the degradation of aromatic and sulphur-bearing amino acids during acid hydrolysis in the presence of carbohydrate (Lugg, 1933, 1938). However, all of these absent amino acids have been recovered in acid hydrolysates of *Penicillium notatum* hyphal walls, with the exception of methionine (Crook & Johnston, 1962) and tryptophan (Applegarth, 1967). Since the carbohydrate content of *P. notatum* does not differ significantly from *P. chrysogenum*, and the conditions of hydrolysis in those studies were essentially the same as used here, it appears unlikely that the amino acids absent from the hydrolysates, with the possible exception of methionine and tryptophan, were initially present in either complex. With respect to the four amino acids found in conidial wall hydrolysates (arginine, lysine, threonine, glycine) but not found in hyphal wall hydrolysates, there is little doubt that they were not part of the hyphal wall complex. Had they been present initially in the hyphal walls, one would expect that their destruction would have occurred during hydrolysis of both conidial and hyphal walls, since both kinds of walls contain approximately the same amount of carbohydrate and were hydrolysed under identical conditions.

The amino acid data for hyphal walls are not in agreement with the values reported for hyphal walls of *Penicillium notatum* by Applegarth (1967) who found threonine, glycine, arginine and lysine as components of hyphal wall hydrolysates, the latter two as major constituents (all absent from hyphal walls in *P. chrysogenum*).

Quantitatively, the most significant difference appears in the total amino acid content of the two types of wall of *Penicillium chrysogenum*. Conidial walls appeared to contain 10 times as much amino acid as hyphal walls. The value of 1.77% for total amino acid is in good agreement with the figure of 2% suggested by Hamilton & Knight (1962) for hyphal walls of the same organism.

Table 3. Recovery of nitrogen in hydrolysate components and in Kjeldahl digests of conidial and hyphal walls of *Penicillium chrysogenum*

<table>
<thead>
<tr>
<th>Calculated from components</th>
<th>Spore wall (µg. N/mg. wall)</th>
<th>Hyphal wall (µg. N/mg. wall)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amino Acid N</td>
<td>24.60</td>
<td>1.97</td>
</tr>
<tr>
<td>Glucosamine N</td>
<td>8.99</td>
<td>14.11</td>
</tr>
<tr>
<td>Total N</td>
<td>33.59</td>
<td>16.08</td>
</tr>
<tr>
<td>Found in Kjeldahl digests of walls</td>
<td>36.57</td>
<td>17.00</td>
</tr>
<tr>
<td>Calculated/found × 100</td>
<td>91.8%</td>
<td>94.6%</td>
</tr>
</tbody>
</table>

When the total contributions of each of the nitrogen-containing components are summed, excellent agreement is obtained with the direct nitrogen analysis of walls (Table 3), 92% of conidial wall nitrogen and 95% of hyphal wall nitrogen can be accounted for as glucosamine and amino acids; the data support the validity of the quantitative method used, and the high recovery of nitrogen supports the contention that certain amino acids were not present in the cell walls.
DISCUSSION

The significance of any analyses of fungal cell walls, particularly in regard to amino acid composition, depends upon the purity of the preparation. The ‘cleanliness’ of wall preparations has been judged by a variety of criteria. Crook & Johnston (1962) considered the isolation of the walls to be complete when all materials which stained intensely with methylene blue had been removed, or, as in the cases of Sporobolomyces roseus and Penicillium notatum, when cytoplasmic particles could not be detected by examination with the electron microscope. Applegarth (1967) washed the disrupted hyphae of P. notatum several times, but still detected some protein contamination of the preparation by electron microscopy. He then treated the material with 0.1% trypsin for 24 hr, after which electron microscopic examination showed a ‘good’ wall preparation. Mahadevan & Tatum (1965) washed the disrupted hyphae of Neurospora crassa with water and increasing concentrations of ethanol and determined the cleanliness of the material by microscopic examination. In the present work, the cell walls were judged pure when ultraviolet absorbing material was no longer released during the series of washings described. None of these criteria, however, is wholly reliable for judging the purity of walls, since adsorbed materials would not be detected. Even if it could be demonstrated unequivocally that the wall preparations were uncontaminated by cytoplasmic material, the preparations might differ from the native complexes. In preparations treated with lytic enzymes after disruption but before hydrolysis, it is difficult to determine whether the enzyme treatment is removing contaminating substances or removing material native to the complex as well or if the enzyme has become adsorbed. An indication that the walls used in the present work were not contaminated by cytoplasmic material is that the preparations lacked several amino acids normally found in proteins. The absence of these amino acids also argues for the absence of cytoplasmic membranes if the assumption is made that fungal membranes are similar to bacterial membranes, which contain all the known amino acids (Weibull & Bergstrom, 1958; Brown, 1961; Yudkin, 1966). The alternative assumption, that fungal membranes are radically different from bacterial membranes and do not contain certain amino acids allows for the possibility of membrane contamination. This appears to be unlikely, since the data would require not only that hyphal membrane be different from conidial membrane, containing even fewer amino acids, but in addition, the amino acids present in the two types of membrane would have to be present in different proportions.

Some insight into the nature of the polymers is provided by the data. In both types of wall, the F 3 fraction is probably a glucan. A similar fraction has been isolated from hyphal walls of Neurospora crassa which was susceptible to the β-1,3-glucanase complex derived from Streptomyces c-3 (Mahadevan & Tatum, 1965). The finding of only glucosamine in the F 4 fractions suggests a chitinous material, as was reported for the F 4 fraction of N. crassa. Such components have also been reported for other fungi (Applegarth, 1967; Chattaway, Holmes & Barlow, 1968; Grisaro et al. 1968). The kinds of polymers in our F 1 and F 2 fractions are considerably in doubt, since these fractions are so complex. No information is available whether these multicomponent fractions represent sets of homopolymers or heteropolymers. Possibly, there might be micelles of one polymer embedded within a matrix of another polymer. Some of the polymers of Penicillium chrysogenum hyphal wall have been degraded enzymically
but enzymic digestion of conidial wall fractions has not been attempted. Intact conidia, however, have been treated in this laboratory with a variety of enzymes in an attempt to produce conidial protoplasts, but unsuccessfully. The intact conidia were resistant to the action of 'Pronase' chitinase, keratinase and the complex of enzymes derived from snail digestive juice. That conidia are refractory to snail enzyme has been corroborated by C. C. Remsen (personal communication). Hyphal walls of Aspergillus phoenicis are extensively digested by a mixture of purified chitinase and B-1, 3-glucanase, but the conidia of this organism are resistant to these enzymes, although they contain both glucose and N-acetylhexosamine in their walls (Bloomfield & Alexander, 1967).

The data in Table 1 suggest the possibility of a lamellar arrangement of wall polymers. Most of the glucose in the conidial walls was released by the second NaOH treatment, rather than the first, and thus required the acid extraction between the NaOH treatments in order to be solubilized. It seems likely, therefore, that the F3 fraction, in which most of the glucose is located, is covered by a base-resistant material which prevents F3 from being solubilized simultaneously with F1. Similar arguments could be advanced for the fact that glucosamine appears in both F2 and F4, rather than F2 only. Preliminary examination of thin sections of conidia by electron microscopy appear to substantiate the presence of a lamellar array. The inability to convert conidia to protoplasts, therefore, may have resulted from the absence of one or more of the enzymes necessary for the sequential hydrolysis of the several major structural polymers of the cell wall.

The differences in distribution and concentration of the sugars and amino acids clearly indicates that conidial and hyphal walls are biochemically distinct.

REFERENCES

**Conidial and hyphal walls**


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