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# Preliminary Experimentation in the Development Of Natural Food Analogues For Culture of Detritivorous Shrimp



# Steven Y. Newell and Jack W. Fell



Technical Bulletin Number 30 July, 1975

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# PRELIMINARY EXPERIMENTATION IN THE DEVELOPMENT OF NATURAL FOOD

# ANALOGUES FOR CULTURE OF DETRITIVOROUS SHRIMP

Steven Y. Newell and Jack W. Fell

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Information Dissemination University of Miami Sea Grant Program 1541 Brescia Coral Gables, Florida 33124

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### ABSTRACT

The fungi are the primary mediators of the microbial degradation of vascular plant debris to particulate detritus in aquatic and estuarine ecosystems. Several estuarine animals of commercial food value including penaeid shrimp are known to be nutritionally dependent, at least in part, on the microbial-detrital complexes resulting from degradation of vascular plant debris. Thus, fungi form a natural part of the diets of these animals.

Recent extensive research has demonstrated that the fungi are very attractive as agents of the manufacture of nutritious microbial protein on an industrial scale. The cost of feed is one of the factors which yet prevents the commercial profitability of aquaculture of penaeid shrimp. If, as seems reasonable, agricultural by-products can be converted inexpensively into fungal materials simulating the naturally eaten microbial-detrital complexes, effective low-priced feeds may be developed. This technical bulletin describes a 3-year preliminary effort at testing of this hypothesis.

Fungal fermentations of agricultural by-products were conducted and attempts were made to optimize fungal protein production. Resultant artificial detrital feeds were tested on a proto-commercial scale in outgrowth of penaeid shrimp. Yields of shrimp ranged from very poor to encouragingly high, considering conditions imposed. Screening was begun of fungal capability to degrade the least expensive types of agricultural by-products, when these were modified to reduce lignin-associated refractoriness. Perhaps the most significant of the conclusions to be reached is that successful development of artificial detrital feeds is not likely to be realized without a major commitment of time, expertise, and money. Suggestions are presented for future research directions, and range of applicability is briefly discussed.

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# PRELIMINARY EXPERIMENTATION IN THE DEVELOPMENT OF NATURAL FOOD ANALOGUES FOR CULTURE OF DETRITIVOROUS SHRIMP

### INTRODUCTION

### Studies of formulation of artificial feed for shrimp

ne unrealized potential of shrimp mariculture as a viable commercial industry

nt from the value of the current shrimo fisheries industry (Roedel 1973). The cost of the feed necessary to bring about the growth of satisfactory, marketable crops of penaeid and Macrobrachium shrimp is one of the major obstacles to the emergence of a profit-producing shrimp mariculture industry (Anderson and Tabb, 1971; Idyll, 1973; Shang, 1974; Webber, 1973). The profitable shrimp culture industry of Japan is dependent upon the high market price of shrimp in that country (Bardach et al 1977) but even in Japan feed casts are presenting presching alardisentations a.... research into artificial feed preparations is underway there (Deshimaru and Shi >, 1972). In the United States, where shrimp prices are considerably lower, profital has eluded those business organizations which have sought to achieve it in sh D farming, and research into formulation of inexpensive, yet effective feeds is or ιf. the steps being taken to overcome this problem (Neal, 1973).

> Results of feed formulation research have varied among researchers, conclusions are difficult to draw. There is disagreement regarding the optimal le of major and minor components; for example, Deshimaru and Shigeno (1972) indica that artificial feeds for <u>Penaeus japonicus</u> should contain at least 60% protein more, while Andrews <u>et al.</u> (1972) concluded that 28-32% protein produces the l growth of penaeids. Shigeno <u>et al.</u> (1972) found a positive correlation of pro content (between 60% and 80%) with feed efficiency, and Balazs <u>et al.</u> (1973) belief that their data demonstrated that the quimal growth is greater of 40%, but Shewbart <u>et al.</u> (1973) found that the optimal protein content lay between

22.5% and 30.5%. Zein-Eldin and McGaffey (1976) found what is probably part of the reason for these apparent conflicts; their work showed that percent protein of over 50 gave the best growth results when one protein source was used, but when a different source was used, 32% protein produced better growth. They suggested that protein quality affected the relationship between shrimp growth and percent protein in the diet. Other feed component requirement levels about which there is seeming disagreement in the literature include aliphatic lipids and their fatty acids, carbohydrates, sterols, glucosamine or chitin, and binding and attracting agents. The clearest conclusion to be drawn from examination of the above and other recent studies of shrimp ration requirements (Andrews and Sick, 1973; Balazs et al., 1975; Deshimaru and Kuroki, 1974a-c, 1975; Forster, 1971, 1973; Forster and Gabbott, 1971; Hysmith et al., 1973; Kanazawa et al., 1970, 1971; Kitabayashi et al., 1971a-c; Meyers and Zein-Eldin, 1973; Shudo et al., 1971; Sick and Andrews, 1974; Sick and Harris, 1975; Venkataramiah et al., 1975; Zein-Eldin and Meyers, 1974) is that the interacting effects of protein, lipid, carbohydrate, and other components in shrimp diets make the optimal formulation of artificial feeds very difficult.

Taub (1973) has noted that one means of finding satisfactory diets for aquacultured animals is to simulate their natural diets. The natural diets of penaeid and <u>Macrobrachium</u> shrimp may include sizable quantities of the plant detritus produced in the estuaries and bordering waters which these shrimp inhabit (Bardach <u>et</u> <u>al.</u>, 1972; Condrey <u>et al.</u>, 1972; Cook and Lindner, 1970; Costello and Allen, 1970; Dall, 1967; Ling, 1969; Odum and Heald, 1972). One approach to determination of an adequate diet for these animals is to simulate this natural plant detritus.

2. The formation of plant detritus in estuaries



Florida estuary, finding that the food web of this shrimp nursery ground (Kutkuhn, ...1966) was based lacely on the production of laces quantities of debris hy managements. This plant debris was found to undergo microbial biodegradation to detrital particles which were higher in protein content than the original material (Fell <u>et al.</u>, in press; Newell, 1975). These microbial-detrital complexes served as a food source for meiofaunal and larger detritivores of the estuary. The pattern of dependence of food ...webs\_proon\_secondary..microbial\_production\_applies\_generally\_to\_gowatic\_ecosystems with relatively large ratios of shoreline length to water-surface area (Mathews and Kowalczewski, 1969).

Recent studies have demonstrated that the phenomenon of protein enrichment of low-protein submerged plant debris is related most directly to the activity of fungi (Bärlocher and Kendrick, 1974; Gessner <u>et al.</u>, 1972; Kaushik and Hynes, 1971; May, 1974; Triska, 1970; Willoughby and Redhead, 1973). Fatty acid enrichment due to <u>fungal colonization of plant debris bas, also been indicated (Schultz and Quing 1973)</u>. It is reasonable: to conclude, therefore, that shrimp in estuarine habitats: derive a portion of their nutrition from the fungi which they ingest in micropial-defrita.

Recognition of the fact that conventional methods of production of protein for human consumption are not meeting the present-day need, and that this situation will continue to worsen, has led to a great deal of research into development of more efficient means of production of high protein material (A. Jones, 1974). Much of this research has dealt with the production of microbial or single-cell protein (SCP), and it is reviewed by Mateles and Tannenbaum (1968), Kihlberg (1972), and Porter (1974a, b). Substrate sources for the fermentation processes involved in the production of SCP have included high-starch-content agricultural products (e.g., cassava) and a wide variety of agricultural and food-industry wastes. Although filamentous fungi are not single-cell organisms, they are generally included among agents of SCP production. Reviews of the use of fungi as producers of protein, food and feed are given by Cooke (1973), Gray (1970), Hesseltine (1968), Litchfield (1968), Solomons (1975), Thatcher (1954), and Worgan (1968, 1973). The principal advantages in the use of filamentous fungi in SCP processes are economic. In submerged culture fermentations, because of their growth form, these fungi can be more easily harvested, by basket centrifugation or other coarse filtration methods, and the resultant product is more easily processed into textured food or feedstuff than those of single-cell organisms (Spicer, 1971, 1973; Worgan, 1973). Perhaps the greatest economic advantage of filamentous fungi lies in their amenability to a less conventional (in the West) type of fermentation, viz. solid state fermentations. This point will be further explored below (Section 15).

Recent reports of filamentous fungal protein production research include (substrate in parentheses): Barnes <u>et al.</u> (1972), Biodeterioration Information Centre of the University of Aston in Birmingham, England (waste paper); Bednarski <u>et al.</u> (1970, 1971), Agricultural University of Olsztyn, Poland (milk whey); Brook <u>et al.</u> (1969), Tropical Products Institute, England (cassava); Chahal and Gray (1970) and Chahal <u>et al.</u> (1972), Punjab Agricultural University, India (wood pulp and rice); Church <u>et al.</u> (1972), North Star Research and Development Institute, Minnesota (corn-canning waste and soybean whey); Imrie (1973, 1975), Sekeri-Pataryas <u>et al.</u> (1973) and Christias <u>et al.</u> (1975), Tate and Lyle Research Center, England, Greece, and Belize, Central America (carob bean extract and citrus waste); Poole and Smith uw 3311Dniversitysion. Aberbeerig counfrantlated wy and educate at al., (1974). Literarysistic of

# -Gudph, Canada (cassava) Rogers=and Spino (1973) - National Environmental Research

Center, Ohio (lignocellulosic wastes); Spicer (1971, 1973), Solomons (1973, 1975), Solomons and Scammell (1974a, b), and Anderson <u>et al.</u> (1975), Lord Rank Research Centre, England (several carbohydrates); Thanh and Simard (1973), Laval University, Canada (domestic sewage); Torev (1973), Bulgaria, where industrial-scale production of polypore mycelium for use as human food ("sausages") is now operating (according to Von Hofsten, 1975); Updegraff (1971), University of Denver, Colorado (waste paper); Von Hofsten and Von Hofsten (1974), University of Uppsala, Sweden (cereals and cereal by-products); Wiener and Rhodes (1974) and Griffin <u>et al.</u> (1974), Northern Regional Research Laboratory, Illinois (feedlot wastes); Worgan (1973; and J. Sci. Fd. Agric., in preparation), National College of Food Technology of the University of Reading, England (food processing wastes). Even waste plastics are being used, with some success, as substrates in the production of filamentous fungal protein (Brnwo et <u>al.</u>, 1974). Perhaps the most concerted effort has been that of the Lord Rank Research Centre, which has progressed to the point of attempting satisfaction of international food-safety regulations with its fungal protein product, having proven the nutritive value and non-toxicity of their fungal protein product for rats, chickens, pigs, calves, and baboons (Duthie, 1975).

## 4. Use of fungal biomass as feed

Many of the above researchers have tested their experimental protein products as feed for laboratory animals (rats, mice, chickens) and positive results have been obtained with regard to growth potential. For example, Reade <u>et al.</u> (1974) reported that a protein efficiency ratio of 2.3 was achieved when their <u>Aspergillus</u> product was fed to rats. Solomons (1973) reported that true protein digestibility (by rats) of one of his filamentous fungal products was virtually 100%. This same product gave growth results (and non-toxicity) equal to that of a casein diet with baboons (Duthie, 1975). Church <u>et al.</u> (1972) reported equal growth rates among rats fed a casein-protein diet and rats fed a fungal <u>(Trichoderma viride)</u> – protein diet. Evidence for the utility of fungi as a feed source has also come as an outgrowth of research intended to pinpoint the causes of feed quality reduction or toxicity by moulding; positive effects on growth were discovered in some farm animals (chickens, sheep, pigs) and laboratory

animals (mice, rats) when specific moulding conditions (physical and chemical fermentation conditions and fungal strains) were used (Chah et al., 1973; Fritz et al.,

1973; Marasas and Smalley, 1972; Richardson <u>et al.</u>, 1967; Sharda <u>et al.</u>, 1971; Thomke, <u>raunas - ac-10.365; blue Doubol: et-al. (1965) and Dector et.al.</u> (1968) achieved noritive growth 17726. ... ice and rats with the mycelial by-product of penicillin production used as results in t for soybean and casein fractions of diets. See Anon. (1971) for a replacement ' of further reports of this kind.

growth-promoting qualities of fungal feeds are not surprising if the itritive quality of fungal mycelium is taken into account. The Reade and ns products mentioned in the foregoing paragraph contained 48% (crude) 1 50% (true, = total amino acids) protein respectively. Although most roteins are low in methionine content relative to the requirements of the which they might be fed, the range of methionine contents of filamentous hes above that of the FAO minimum requirement level for humans et al., 1975; Rhodes et al., 1961). Nonetheless, as Church et al. (1972) and 1972) point out, methionine addition is commercially feasible since this is now inexpensive and easily obtainable. Furthermore, Kanazawa et al. Sick and Anderson (1974) have found that soybean meal, which is / low in methionine content," served better as a protein source for penaeid n some animal proteins. Other than the sulfhydryl amino acids, fungal general contain nutritionally satisfactory levels of essential amino acids, a wide range of specific amino acid contents. This is true even among the same genus (Rhodes et al., 1961). Therefore, if one were to search for a rce having a given amino acid balance, it is likely that a species or strain build be found possessing that quality.

ps most notably important in this regard is the tractability of the protein fungi. Levi and Cowling (1969), Litchfield <u>et al</u>. (1963), Merrill and Cowling

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(1969) demonstrated that fungi were able to adapt their nitrogen and protein contents to the C/N ratios of their substrates. Their reports, along with that of Graham (1971), show that the increase in nitrogen reflects an increase in protein content, not in glucosamine (or chitin) or cellular ammonia. Alteration of other physical and chemical conditions of culture can also be used to adjust protein and amino acid content, including methionine content (Chebotarev and Zemlyanukhin, 1973; Florentino and Broquist, 1974; Graham, 1971; Pinto, 1963; Solomons, 1973; Tkachenko <u>et</u> <u>al.</u>, 1972; Verona <u>et al.</u>, 1973). In addition, the chitin content of cell walls of higher filamentous fungi varies greatly from species to species, and the chitin percentage of mycelial biomass varies greatly with age of culture (Blumenthal and Roseman, 1957). The wide range of contents of nutritively valuable components of fungal mycelium together with the manageability of the relative amounts of these components in a given species, validate Gray's (1970) "optimistic view that if one searches far enough and widely enough, he can often find a fungus to perform the particular synthesis { or serve the purpose } in which he is interested."

Three studies of the use of filamentous fungi which relate directly to the potential use of fungal feeds for shrimp are those of Bärlocher and Kendrick (1973, 1975), Joshi (1974), and Nikolei (1961). These are reports of the use of fungal tissue in the culture of arthropods which feed naturally on dead, fungal-decayed plant material. The Bärlocher and Kendrick studies were intended as sequels to the studies of Kaushik and Hynes (1971) and Triska (1970) (Section 2) who had shown that the fungi were responsible for the major portion of the protein increase in decaying plant debris, and that detritus-feeding crustaceans definitely preferred fungal-invaded material over sterile material. Bärlocher and Kendrick fed leaves with a minimal microbial population, leaves richly colonized by microbes, and fungal mycelia (10 species), separately to cultures of a detritivorous gammarid amphipod. Some of the fungi proved to be excellent growth-promoting food, better than the leaf material,

while other fungi produced little or no growth.

Joshi fed fungal meals of differing types to a mycophagous thysanopteran; he found that the length of the larval period and adult fecundity were markedly affected by the species of fungus used in preparation of the meals.

Nikolei found similar results in his search for suitable fungal tissue to serve as a culture medium for three species of cecidomyiid flies. He tested 67 species of fungi and seven strains of <u>Trichoderma viride</u>. Fifty of the species were unsuitable for culture of the fly larvae and of the seventeen which were suitable, only four were suitable for all three fly species. There was considerable variability among the <u>T</u>. <u>viride</u> strains in nutritive capacity which ranged from highly positive to distinctly negative. Also, as Bärlocher and Kendrick had found or suspected, culture conditions and age of the fungi had marked effects on their growth-producing potential. Again, Gray's (1970) "if one searches far enough ..." viewpoint seems to be borne out.

Although the present authors know of no published reports of the feeding of fungal mycelia to postlarval or juvenile commercially valuable shrimp, the following reports indicate its plausibility. R. R. Jones (1974) has demonstrated that, as in the case of other detritivorous crustacea (Fenchel, 1972; Hargrave, 1970), natural detrital materials are nutritively utilized by young juveniles of <u>Penaeus aztecus</u> with high protein assimilation efficiency. Even the cell walls of the fungi probably serve as sources of shrimp nutrition. The cell walls of the higher filamentous fungi are chitinous (LéJohn, 1971), and Hood and Meyers (1974; and unpublished communication) have shown that penaeid shrimp have, in addition to a hepatopancreal chitinase, a bacterial gut flora which is 85% chitinolytic and which increases in size with addition of chitin to the diet of the shrimp. Minami <u>et al.</u> (1972) have shown that the size of the chitinolytic bacterial population of fish guts is positively correlated with their efficiency of utilization of petro-yeast. The requirement of penaeids for a chitin component in their diets has been examined, but disagreement concerning the nature

of this requirement exists (compare Deshimaru and Kuroki, 1974b; Kanazawa <u>et al.</u>, 1970; Kitabayashi <u>et al.</u>, 1971a).

The following reports are also of interest in this connection. Ewald (1965) used marine yeast (a <u>Rhodotorula</u> species) as a component of the food which he used in the early successful work with raising of larval <u>Penaeus duorarum</u>. Forster (1973) found that the nitrogen in a yeast-protein component of a palaemonid shrimp ration was assimilated at 95% efficiency, and Andrews and Sick (1973) found that yeast-protein labeled with carbon 14 was incorporated at a high rate into penaeid shrimp muscle tissue. Meyers <u>et al.</u>, (1970) found that young freshwater crawfish grew "encouragingly" when fed a diet consisting of 39% spray-dried yeast. In a very important personal communication, Dr. E. B. G. Jones (Portsmouth Polytechnic, England) reports that he and his colleagues have found that commercially culturable shrimp <u>will ingest</u> fungal mycelia, and that "reasonably good" yields have resulted.

# MATERIALS, METHODS, AND RESULTS

## 5. History of an artificial detrital feed project

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As an adjunct project, we (unpublished data) conducted a microbiological and gross biochemical survey of the artificial detrital materials from the shrimp tanks over a period of four weeks. Litter-bag sampling methods similar to those of Newell (1975) were used. One major deviation from those methods was the added emphasis placed on results from incubation of pieces of the materials in dishes of sterile seawater (15 o/oo) with antibiotics (penicillin and streptomycin, 0.05%).

The survey showed that, although distinct fungal florae had inhabited the different detrital material (Table 1)\*, nitrogen enrichment of the materials did not take place. During the four weeks of sampling fungal colonization increased as bacterial colonization decreased, and the materials lost about half of their original nitrogen content. The biodegradation phenomena which Caillouet and Tabb had hoped for did not take place in their tanks. The major reasons for this were probably: (1) that significant increase in nitrogen content of lignocellulosic materials due to fungal biodegradation requires more time than was allowed; (2) that the conditions requisite for permitting the fungal biodegradatory process were not supplied; (3) little or no natural inoculum was present (note { Table 1} the marked differences in fungal occurrence patterns in the tanks as opposed to the adjacent mangrove pond); and (4) unavailability of macronutrients and buildup of fungal growth inhibitors may well have been brought about as a consequence of the standing condition of the water of the tanks. Yield of shrimp in the experiment, rather than being related to extent of

<sup>\*</sup> The following are points of note from Table I. The bagasse had the heaviest fungal inoculum at the outset. Further colonization was most extensive on bagasse in the tanks, but the difference in extent of colonization was less marked in the mangrove channel. Some fungi present initially on the substrates were not reduced in frequency of occurrence by submergence (e.g., <u>Trichoderma viride</u>, <u>Syncephalastrum</u> <u>racemosum</u>) while others were eliminated (<u>Chaetomium sp.</u>), or reduced (<u>Alternaria</u> <u>sp.</u>). Some fungi colonized more extensively in the tanks (<u>Stachybotrys</u> sp.), while others were equally frequent in the mangrove channel (<u>Trichoderma viride</u>, <u>Zythia</u> sp. on straw). The obligate marine fungi (<u>Lulworthia</u> sp., <u>Zalerion</u> varium, <u>Thraustochytrium pachydermum</u>) colonized only in the mangrove channel.

Table 1. Frequency (%) of occurrence for the prevalent fungi which colonized materials used as artificial detrital feeds in mariculture of penaet shrimp.
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	HM	EAT BRAN	t-a l	3¥I	HEAT ST	RAW		BAGASSI	real
	PBa	TK	MC	PB	TK	MC	PB	TΚ	MC
no fungi	100	71	4	89	34	13	44	'n	2
diatoms (heterotrophy) <sup>b</sup>	0	28	0	0	20	0	0	0	o
sterile mycelium	o	25	25	0	33	27	Q	'n	5
Ryphomycetes									
<u>Alternaria</u> sp.	0	0	0	11	e.	0	0	ς'n	0
Ascotricha chartarum	0	0	0	0	0	0	0	7	0
Aspergillus spp.	0	0	0	0	0	0	0	80	11
Cephalosporium spp.	0	0	15	0	0	12	Q	57	65
Cirrenalia pseudomacro- cephala	0	0	4	o	0	, N	o	0	Q
Cladosporium sp.	0	2	6	0	0	0	0	0	83
Colletotrichum sp.	0	0	4	0	0	0	o	0	0
Culcitaina achraspora	0	0	0	o	0	ო	0	0	0
Dendryphiella arenaria	0	0	o	0	0	0	0	0	ø
Fusarium sp.	0	0	0	0	0	0	0	7	0

H

Table 1 (continued)

	WHE	AT BRAN		5	HEAT STH	RAW		BAGASSE	
	PB <sup>a</sup>	Ħ	MC	PB.	¥	MC	BI	ä	Щ
Paecilomyces sp.	0	0	4	0	0	0	0	9	0
Penicillium spp.	0	0	7	0	0	0	0	21	29
Stachybotrys sp.	0	0	0	o	0	e	0	42	13
Irichoderma viride	0	0	0	0	0	0	9	23	22
Zalerion varium	0	0	o	0	0	٢	0	0	'n
Coelomycetes									
Z <u>ythia</u> spp.	0	0	11	0	12	12	ę	17	Ń
Ascomycetes									
<u>Chaetomium</u> sp.	0	0	0	0	0	0	22	¢	0
Lulworthia sp.	0	0	13	o	0	ഹ്	0	0	Ś
Zygomycetes									
Syncephalastrum racemosum	0	0	0	0	0	0	9	10	Ś
Chytridiaceous fungi									
Thraustochytrium pachydermum <sup>c</sup>	I	0	33	I	0	27	I	0	61
Phlyctochytrium mangrovii <sup>c</sup>	1	0	10	I	0	11	I	0	16

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	IJ	EAT BRAN		12	HEAT STI	RAW		BAGASSI	r-11
	PB <sup>a</sup>	TK	WC	PB	TK	MC	BB	Я	MC
bactería <sup>d</sup>	I	0.10	0.16	۱	< 0.01	0,09	I	0.04	0.05
nze	2.6	1.3	1.8	0.4	0.1	0.1	0.4	0.2	0.4
number of observations	18	126	55	18	126	63	18	126	60

- shrimp tanks; MC=sampling after placing in litter bags in a mangrove lined estuarine channel adjacent <sup>a</sup> PB=sampling prior to placement in litter bags; TK=sampled periodically after placing in litter bags in to the shrimp tanks.
- <sup>b</sup> Includes only those observations made on pieces of the material which were washed, surface-sterilized, rinsed, and plated on cornneal agar.
- v. 1. -/ UU BEAWATER WITH 0.05 % penicillin and streptomycin, and baited with boiled blades of drop-seed grass (Sporobolus virginicus). <sup>c</sup> Includes only those observations made on pieces of the material which were washed and incubated in dishes of 15 °/oo seawater with 0.05 %
- <sup>d</sup> The figures given are the ratios of final number of viable elements to initial numbers of viable elements per piece of material (determined by membrane (0.45 µm) filtration of dilutions of the wash water, and plating of the membranes on a glucose-peptone-yeast extract agar).
- <sup>e</sup> Nitrogen percentage at final sampling. Comparison of the PB samples with TK and MC samples show change in NX from original.

biodegradation, was positively correlated with the original nitrogen content of the materials; shrimp fed wheat bran (N% = 2.7) were on the average over twice as large as those fed the nearly purely lignocellulosic straw and bagasse (N% = 0.4).

### 6. Culture-screening on wheat bran and bagasse

### Background.

The next logical step in the development of artificial detrital feeds was an acceleration of the natural biodegradation process. This we hoped to accomplish by moving the process from the shrimp tanks into separate containers in which the biodegradation could be conducted as controlled, submerged-culture, fungal fermentations. We planned our fermentation experimentation for use of diluted (15 o/oo) seawater because: (1) penaeid culture is most economically accomplished as a coastal process; (2) fresh water is one of the earth's threatened resources; (3) Gray et al. (1963) had shown that use of seawater in fungal fermentations could produce enhancement of protein production; (4) the estuarine-adapted fungi form a portion of the natural shrimp diet (Section 2); and (5) both terrestrial and obligately marine fungi in general can proliferate in partial seawater (Jones et al., 1971).

Materials and methods.

We began by conducting a culture screen of 29 isolates of fungi and 5 mixed cultures of fungi for their ability to increase the proteinaceous material in wheat bran and bagasse. A diversity of fungal species was used (Table 2), including fungi restricted to marine-estuarine environments (e.g. <u>Lulworthia</u> spp.), fungi isolated from the plant materials during the Caillouet-Tabb shrimp-feeding experiment (e.g. <u>Stachybotrys</u> sp.), and other fungi recently isolated from naturally decaying plant materials in local estuaries, some of which had been indicated as potentially efficient protein producers (e.g. Trichoderma viride by Church <u>et al.</u>, 1972).

Shake flask fermentations were performed as prescribed by Solomons (1969) and by adapting the methods of Sguros <u>et al.</u> (1962). A cylinder ( $2 \text{ cm}^2 \text{ diam.}$ ) was

removed (sterile #14 cork borer) from the growing edge of a colony of each of the test fungi on cornmeal (Difco) agar (CMA) made with 15 o/oo seawater. The cylinder was placed in the bottom of a presterilized Monel Metal Waring microblender in 25 ml of sterile 15 o/oo seawater. and homogenized for 60 seconds at low speed. One ml of the

homogenate was added, in a 125 ml DeLong flask, to 25 ml of the inoculum-growth medium: glucose - 1%, NH<sub> $\mu$ </sub>NO<sub>3</sub> - 0.24%, KH<sub>2</sub>PO<sub> $\mu$ </sub> - 0.006%, yeast extract (Difco) -0.01%, cellulose (powdered Whatman filter paper) - 0.01%, pectin - 0.01%, in 15 o/oo seawater. The flasks were capped with Belco metal closures, sealed around the lower edges of the caps with clear tape (to inhibit contamination), and incubated at 28°C on a temperature-controlled reciprocating shaker (70 cycles/min. through 8 cm distance). These inoculum-growth flasks were harvested at approximately the early stationary phase (3 to 9 days). As was the case with Wang et al. (1974), only subjective determinations of growth phase were made due to the difficulties inherent in determining quantities of mycelia mixed with solid substrate residues (Calam, 1969b). The sieve-separation method of Borzani et al. (1972) was not attempted. The entire content of each flask was homogenized for 60 seconds, and 2 ml of homogenate was added, in each of two 125 ml final culture-screen flasks, to 25 ml of the following media: one of 5% ground bagasse (40-mesh) and one of 5% ground wheat bran, with the remainder of the media the same as that of the inoculum-growth medium, with the omission of the cellulose and pectin. These final flasks were incubated as were the inoculum-growth flasks.

Ideally, the inocula would have been washed and their dry weights standardized. This was avoided for the following reasons: impracticability in view of limited time and personnel; severe increase of contamination risk; ineffectiveness of this method of standardization of inoculum potential. Our work (Table 5) has shown that equal times of homogenization produce large variations among different fungi in numbers of viable mycelial elements per unit dry weight. This can have a marked effect on inoculum potential (e.g. Graham, 1971). The mycelium exhaustion steps of the inoculum preparation of Sguros <u>et al.</u> (1962) were not used in the present experiment, as this would certainly not be a part of production-scale fermentation processes, in which optimization of vigor of inoculum is desired.

The work of Sguros et al. (1962) and Ward and Colotelo (1960) on systematically divergent fungi both showed that amounts of inoculum of dry weight greater than 2 mg (into 25 ml of medium) do not have marked effects on final yield of mycelium. We calculated from the papers of Sguros et al. (listed in Sguros, 1973) and Meyers (reviewed in Meyers, 1971) that 2 ml of homogenate produced as above would contain at least 2 mg of mycelium. Upon inoculation of each culture screen flask, a second, control 2 ml of homogenate was vacuum-filtered on a tared glass filter pad (Whatman GF/A), washed with 50 ml deionized water, and dried at  $55^{\circ}$ C. The range of inoculum dry weights was 2.3 mg (Stachybotrys kampalensis SC76) to 11.7 mg (Pestalotia sp. SC 38). In order to determine the effect of unequal inocula, a regression was performed, of final nitrogen percentage of fermented wheat bran on inoculum weight. Significance was detected (p = 0.001) with a slope of - 0.11 and the coefficient of determination  $(r^2) = 0.25$ ; those fungi which produced heavier inocula tended to produce lower final nitrogen percents in fermented wheat bran, and 25% of the variation in percent nitrogen was due to inoculum size. This was probably primarily a function of the positive correlation between inoculum size and growth rate: the fastest-growing forms produced larger inocula, and were likely to be harvested at a point in their growth cycles beyond early stationary phase.

Harvesting of fermented product was done at approximately early stationary growth phase by coarse (42  $\mu$  m), double-cloth filtration, with tap water washing until soluble nitrogen was removed (2 equal volumes of tap water, with intervening through resuspensions). The products were then dried at 55°C and ground through 60-mesh Wiley mill screen. Nitrogen content was analyzed using a Perkin-Elmer Model 240

Elemental Analyzer, and protein figures given are for crude protein (N x 6.25).

Further screens of fungal isolates for their capacity to convert wheat bran into fungal protein were conducted using a second standard set of fermentation conditions raising the total number of isolates screened with wheat bran to 52. The medium used in the second set of wheat bran screens was: wheat bran, unground - 5%,  $NH_4NO_3 = 0.86\%$ ,  $KH_2PO_4 = 0.09\%$ , yeast extract = 0.01\%, in 15 o/oo seawater. Inoculation of 50 ml of medium in 250 ml Ehrlenmeyer flasks was by addition of 2 ml of CMAA-culture homogeneate-asign the first wheat bran screen (inoculum growth flasks were not used). Incubation, harvesting, and analysis were the same as in the first wheat bran screen.

Results.

Results of the culture screens are given in Tables 2 and 3 and Figure I. The figures are for the solid end products of the fermentations, fermented plant particles plus fungal mycelium. Although only one of the wheat bran products was lower in percent protein than the original material, the range of crude protein increments was from -46.5% (Syncephalastrum racemosum, Table 2) to +50.6% (Myrothecium sp., Table 3). The medium used in the first bran culture screen was 3.6 times lower in inorganic nitrogen than the medium used in the later culture screen. Probably as a partial consequence, there were more losses in crude protein observed in the first wheat bran screen than in the second (Fig. 1). Bagasse conversions (Table 2) were much poorer than with wheat bran. Although there were some high increments in

Table 2. Results of the first culture screening intended to detect strains of fungi optimally able to convert wheat bran and bagasse into mycelial biomass.<sup>a</sup>

FUNGAL STRAINS <sup>b</sup>	WHEAT	BRAN	BAC	GASSE
	<u>CP%</u> C	<u>∆CP%<sup>d</sup></u>	CP%	∆CP%
Lulworthia sp. SC73	28,9	+16.9	2.9	-12.8
SC73 & SC51	27.9	+15.4	2.9	-15,3
Leptosphaeria maritima RZ312	25.8	+14.5	2.7	-15.3
Leptosphaeria discors SC71	25.4	+ 3.3	2.9	-13.0
Zalerion varium SC21	24.1	+ 2.9	2.9	-12.8
Zalerion varium RZ264	26.4	+ 2.2	3.1	- 7.9
Trichoderma viride SC51	24.8	+ 2.1	3.3	- 5.2
Drechslera hawaiiensis RZ17	24.5	+ 0.1	<sup>f</sup>	
Lulworthia medusa V biscaynia RZ 281	26.1	- 0.1	5.3	+53.7
Lulworthia grandispora RZ389	24.4	- 1.5		
Stachybotrys kampalensis SC76	23.9	- 2.1		*
Zalerion varium RZ394	23.8	- 3,5	3.3	- 0.5
<u>Culcitalna</u> achraspora SC24	23.0	- 4.8		
Corollospora maritima SC67	23.4	- 6.0	4.0	+14.5
Stachybotrys sp. SC52	25.3	- 7.1	4.6	+22.2
Culcitalna achraspora RZ387	22.4	- 8,9	3.8	+15.5
Leptosphaeria albopunctata SC70	22.1	-12.2	3.6	+ 4.2
Cirrenalia pseudomacrocephala RZ280	22.8	-12,4	2.8	-17.0
SC47 & SC70	23.0	-13.8	3.5	+ 2.2
<u>Pestalotia</u> sp. SC38	20.4	-14.8	4.0	+20.7
Zalerion varium RZ393	21.7	-15.8	3.2	- 6.2
<u>Clavariopsis</u> <u>bulbosa</u> SC66	22.4	-17,1	5.5	+45.0
Corollospora maritima SC68	22.6	-17.2	6.3	+68.9
Ascotricha chartarum SC44	20.0	-18.4	2.5	-26.1
SC73 & SC62	21.8	-19.3	2.8	-19.0
Hyalostachybotrys sp. SC74	18.7	-24.9	4.7	+40.6
<u>Nia vibrissa</u> SC72	16.9	-25.6	3.6	+ 8.6
SC47 & RZ387	18.9	-28.9	2.6	-21.4
<u>Hyalostachybotrys</u> sp. SC62	19.9	-29.4	3.5	- 1.0

FUNGAL STRAINS <sup>b</sup>	WHEAT	BRAN	BAG	ASSE
	CP% <sup>C</sup>	$\Delta CP \%^{d}$	<u>CP%</u>	ΔCP%
SC47 & RZ393	18.4	-32.5	4.4	+28.8
Robillarda sp. SC75	17.3	-33.5	2.1	-34.7
SC47 & SC73	17.3	-37.4	3.3	- 4.7
<u>Syncephalastrum</u> <u>racemosum</u> SC47	15.1	-46.5	2.2	-34.7
<u>Dendryphiella</u> salina SC77	<b>-</b> -		2.8	-17.0
Processed control <sup>e</sup>	14.3	-44.4	2.6	-22.9
Untreated material	16.7		2.8	

<sup>a</sup> Medium used: 5% wheat bran or bagasse (ground through a 40-mesh Wiley mill screen), 0.24% NH<sub>2</sub>NO<sub>3</sub>, 0.006% KH<sub>2</sub>PO<sub>4</sub>, 0.01% yeast extract, in 25 ml. 15 o/oo seawater in 125 ml DeLong flasks.

<sup>b</sup> With culture collection accession numbers.

- <sup>c</sup> Percent crude protein (= percent nitrogen x 6.25) of fermented product (wheat bran and mycelium).
- <sup>d</sup> Crude protein of final product/crude protein of original material, x 100.
- <sup>e</sup> Flasks of medium taken through sterilization, fermentation and harvesting process, but not inoculated.

f Not determined.

Table 3. Results of the second set of culture screens intended to detect strains of fungi optimally able to convert wheat bran into mycelial biomass.<sup>a</sup>

FUNGAL STRAIN <sup>b</sup>	WHE	AT BRAN
	<u>CP%</u> <sup>C</sup>	$\Delta CP X^d$
Myrothacium sp. SC87	34.3	+50.6
Septonema secedens SC104	29.8	+40.6
Curvularia sp. SC86	33.4	+37.9
Zythia sp. SC20	32.5	+35.9
Chaetomium sp. SC97	31.7	+34.8
Scopulariopsis (?) sp. SC8	31.0	+20.3
Aspergillus terreus SCU14	31.0	+19.6
Lulworthia sp. SC73	29.9	+19.3
<u>Melanospora</u> sp. SC94	27.6	+17.9
<u>Curvularia tuberculata</u> SC91	26.3	+10.1
Aspergillus niger SC90	27,7	+ 7.7
Cephalosporium sp. SC1	28.3	- 0.3
<u>Cladosporium</u> <u>cladosporioides</u> SC 103	22.9	- 1.1
Cladosporium sp. SC102	22.0	- 2.3
Geotrichum candidum SC117	26.4	- 2.3
<u>Alternaria longissima</u> SC122	24.4	- 5.7
Botryosphaeria sp. RZ 254	25.5	- 7.3
Zythia sp. SC7	21.2	-11.8
Dendryphiella salina SC77	26.4	-12.0
<u>Blakeslea trispora</u> SC88	20.6	-15.3
Sporormia sp. SC100	23.7	-15.9
<u>Aspergillus</u> amstelodami SC43	21.7	-20.9
Trichoderma viride SC51	33.1	<sup>e</sup>
<u>Pestalotia</u> sp. SC38	32.9	
<u>Pestalotia</u> sp. SC89	32.5	

Table 3. (continued)

- <sup>a</sup> Medium used: 5% wheat bran, 0.86% NH4N03, 0.09% KH P024, 0.01% yeast extract, in 50 ml 15 o/oo seawater, in 250 ml Ehrlenmeyer flasks.
- <sup>b</sup> With culture collection accession numbers.
- <sup>c</sup> Percent crude protein (= percent nitrogen x 6.25) of fermented product (wheat bran + mycelium).
- <sup>d</sup> Crude protein of final product/crude protein of original material, x 100.
- e Not determined.



Distribution of screened fungi by size class of: A. nitrogen content (N%) of fermented wheat bran; B. percent crude protein change ( $\Delta CP\%$ ) of fermented wheat bran. See Tables 2 and 3 and text for further details. Figure 1.

percent crude protein (<u>Lulworthia medusa</u> var. <u>biscaynia</u>, 53.7%, and <u>Corollospora</u> <u>maritima</u>, 68.9%), the highest final crude protein content of the end products was only 6.3%, considerably lower than the requirement of penaeid shrimp (Section I).

# 7. Production of experimental feeds

Background.

Before the crude protein analyses of the first culture screen could be completed, an opportunity arose for preliminary testing of fungal-fermented products as feed for pink shrimp. Three fungi from the culture screen were used to ferment wheat bran in 5 gallon glass carboys, using the same conditions as in the first culture <u>screen</u>. The products were backested as before, dried at 50°C, ground (brough a.40mesh Wiley mill screen, and fed to postlarval (average initial fresh weight - 0.11 gm) pink shrimp in the concrete tanks described above. It was unfortunate that the crude protein analyses from the first culture screen were not available during the design of this experiment, for the three fungi chosen were among the poorest performers of the screen (<u>Syncephalastrum racemosum</u>, <u>Nia vibrissa</u>, and <u>Dendryphiella salina</u>). Maximum final crude protein content from the carboy fermentations was 18.6%, corresponding to a net loss of 57.0% crude protein. Shrimp yield from all three fungal feeds was 1.6 times or more lower than that from unfermented bran, though all three fungal feeds gave average shrimp yields 7-8 times higher than unfed controls.

Materials and methods.

During the fall of 1973, we obtained a 14-liter laboratory fermentor (Microferm, New Brunswick Scientific Co.), with dissolved oxygen and pH controls, for the purpose of feed production for the 1974 shrimp-growing season. Only the warmer months when temperature does not fall below 20<sup>o</sup>C are suitable (e.g. Kilgen and Harris, 1975). We completed analysis of our culture screens, and identified those isolates apparently best able to convert wheat bran into a more highly proteinaceous product. Fermentor runs were planned using these fungi in order to determine how closely scaled-up
fermentation processes would correspond to the flask-level fermentations, and to produce feed material for shrimp-feeding experimentation in 1974. We soon encountered the difficulty discussed by Solomons (1969), Underkofler <u>et al.</u> (1947), and Weismann (1970) with fermentation of particulate substrates: contamination due to the resistance to conventional steam sterilization of spore-forming bacteria, when these are protected by starch or protein components of particulate media in large volume. Rather than solve the problem by purchase of a stirring autoclave, as Weismann (1970) suggests, or by using in-place steam sterilization (impractical in our case), we found that slight modification of the autoclave and <u>separate</u> autoclave sterilization of all fermentation components (bran, seawater and inorganic nutrients, apparatus) was effective in preventing bacterial contamination. The autoclave modification involved venting from near the bottom rather than the top, permitting saturated steam to displace air (Ernst, 1968).

Because of the potential contamination problem, we used fast-growing (and thus contamination-resistant) fungi as fermenting agents in our production of test feeds. Other considerations here were the necessity to produce large quantities rapidly, and to reduce the projected cost of large scale operation (which reduction in length of the reduce the projected cost of large scale operation (which reduction in length of the reduction in length of the reduce the projected cost of large scale operation (which reduction in length of the reduction in length of the

lum was produced /ere performed as arvested at early seconds as in the ame medium in a .flasks.had.hem / 60 ml aliquots) The fermentation Calam (1969a, b), Rowley and Bull (1973), and Solomons (1969). Inoc as follows: 250 ml Ehrlenmeyer-flask fermentations of wheat bran in the second set of wheat bran culture screens. These were stationary phase, the entire 50 ml contents homogenized for 60 culture screens, and the entire homogenate added to 450 ml of the 2000 ml Fernhard flash. This was incudanted as the 250 ml stat harvested at early stationary phase, homogenized (in approximate and added aseptically to the sterilized medium in the fermentor. was then immediately begun (agitation, sparging, pH, foam, and temperature controls

adjusted and set into operation). Settings and conditions for individual rans are reasonable in Table 4. Foam control was by automatic addition of polypropylene glycol P2000. Except when difficulties occurred with probe-sensitivity fluctuation (not shown in Table 4), only a few milliliters of the antifoam agent were added to the fermentation. All the fermentation runs were temperature-controlled at 28°C. All runs were batch fermentations; we attempted to terminate all runs in early stationary phase, as

indicated by decline in oxygen uptake and rise in pH (all runs were with ammonium ion as the inorganic nitrogen source; see Prokop and Stros, 1974). Harvesting and analytical methods were similar, though on a larger scale, to those used in the culture screens. In addition, solids which passed the double-layer,  $42 \mu$  m nylon-cloth filters were collected by continuous centrifugation in a Sharples TI Super Centrifuge which was obtained at mid-year, 1974.

Results.

The four fungi which were used and the results of fermentor runs, the products of which were used in subsequent shrimp-feeding experiments, are given in Table 4. Range of content of crude protein for the four fungal products was from 25% to 32%, and increment in crude protein from -31% to +25%.

## 8. Attempts to optimize crude protein production

Materials and methods.

During summer, 1974, we performed four flask-scale experiments designed to determine the extent to which wheat bran fermentation results could be improved by manipulation of fermentation conditions. The experimental design was a 3-way, 3-level, 2-replicate, factorial analysis of variance (ANOVA), and each experiment tested the response of a different one of the fungi from the bran culture screens. Statistical methodology used in this experiment and the others of this communication

was derived from Stege: (1956); Shedebor and Cochran (1967); Sokat and Konif (1969),

Conditions and results of fermentations conducted in a 14-liter laboratory fermentor, the products of which were used as test feeds for shrimp in the F74 series (Section 9). Table 4.

t x .9.2 A	აი +	+25	¥	£	a I	<b>11</b>	+13	+ <b>1</b> 1	<b>8</b>	Ŷ	۴ ۲	7	-31	-27	-25
r 7 niejory eburd	28	32	31	32	28	32	31	31	31	25	26	27	26	27	26
A Dry Weight Zh	37	33	41	44	e'	40	38	39	<b>0</b> †	37	40	39	5	54	51
Burstion, hrs. <sup>8</sup>	94	69	73	115	69	96	52	98	92	11	33	37	36	32	33
f	н	2	2	2	2	7	7	7	7	7	2	7	4	7	7
ુખ્રસ	300	500	500	500	500	500	500	500	500	500	500	700	700	600	600
b <b>. 1a , beed , as B</b>	¥	100	120	23	139	0	717	o	0	0	42	62	63	<b>90</b>	10
.ат, <u>ях</u> ,	72	72	72	96	85	120	96	120	72	72	52	96	96	48	48
Inoculum, <sup>1</sup> ts.	0.5	0.5	0.5	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
oo/ viiniis	15	15	15	15	15	15	15	15	15	15	15	15	ŝ	ŝ	ŝ
E ON 7HINZ	0.86	0.86	0.65	0.65	0.65	0.65	0.65	0.65	0.65	0.65	0.65	0.65	0.54 <sup>n</sup>	0.34	0.34
а Втал Втал Втал	ŝ	S	'n	Ś	ŝ	ŝ	ŝ	5	5	Ś	Ś	ŝ	ø	2	7
suisro? Isgnuf	SC51	SC51	SC51	SC87	SC51	SC87	SC51	SC87	SC87	SC86	SC38	SC38	<b>S</b> C51	SC87	SC87
Run Number	19	20	21	22	24	25	26	27	28	29	31	32	34	35	36

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Table 4. (continued)

<sup>a</sup> Fungal strain used, given as culture collection accession numbers. SC51-Trichoderma viride; SC87-Myrothecium sp.; SC86-Curvularia sp.;

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- <sup>b</sup> The percentage of wheat bran used, w/v. Wheat bran was except for runs 35 and 36, for which it was pre-ground Wiley mill screen. Runs 24-36 employed 10.5 liters of 19-24, 11 liters.
- <sup>c</sup> The age at which inoculum cultures were homogenized and fermentation vessel.
- <sup>d</sup> The amount of 2N KOH utilized by the automatic pH contr maintaining the pH above (not at) 4.0.
- <sup>e</sup> The revolutions per minute of the fermentor impeller.
- f The air flow into the fermentation vessel.
- <sup>g</sup> The length of time between inoculation and harvest of t
- <sup>h</sup> Final dry weight of product (mycelium and residual bran weight of bran, x 100.
- <sup>i</sup> Percent nitrogen of product x 6.25.
- <sup>j</sup> Crude protein of product/initial crude protein of bran,
- k pH maintained at 4.5 throughout fermentation.
- <sup>m</sup> Data lost.
- <sup>n</sup>  $(NH_4)_2 SO_4$ .

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and Zar (1974). Two fast-growing fungi were used (Trichoderma viride SC 51 and Myrothecium sp. SC 87) and two slower-growing, marine fungi (Lulworthia sp. SC 73 and Leptosphaeria maritima RZ 312). The three treatments with their levels were: salinity, 5 0/00, 15 0/00, 30 0/00; bran concentration (w/v), 2%, 5%, 8%; type of inorganic nitrogen source (w/v), high  $NH_4N0_3$ , low  $NH_4N0_3$ , and low  $(NH_4)_2 SO_4$ . The levels of inorganic nitrogen salt used provided an equivalent carbon/ammonium ion ratio in the low  $NH_4N0_3$  and  $(NH_4)_2 S0_4$  flasks (C/NH<sub>4</sub> - N = 28) and one-half that ratio for the high  $NH_{ij}NO_3$  flasks. The resultant percentages of the nitrogen sources were, for high  $NH_4NO_3$ : 0.34%, 0.86%, and 1.38%; for low  $NH_4NO_3$ : 0.17%, 0.43%, and 0.69%; for  $(NH_4)_2 SO_4$ : 0.14%, 0.34%, and 0.54%. The C/NH4-N ratio was used for standardization, rather than C/N, because of the probability that the fungi would take up the ammonium ion in preference to the nitrate ion (Nicholas, 1965) and the fact that the ammonium was probably present in large excess of requirement for maximal theoretical nitrogen content of mycelium and maximal theoretical conversion of bran (Section 6). Methods of preparation, incubation, and analysis were similar to those of the first bran culture screen, with two important exceptions: (I) inoculum level was lower than in the culture screen (Table 5; 2 ml of each homogenate were added to each 125 ml experimental flask) and the inoculum for SC 73 was less than 2 mg/flask; (2) all fermentations were terminated at 7 days, rather than at subjectively determined early stationary phase.

Due to the large numbers of flasks having to be harvested at the same time, all flasks of a given experiment were removed from the incubator at the same time, and stored at 2°C. They were then harvested on a random basis over a period of 5 days. <u>A sequence of the product of the product of the period of the store o</u>

## Table 5. Inoculum data for the $3 \ge 3 \ge 3$ ANOVA experiments designed to determine how the wheat bran fermentations might be optimized

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)L. b	<u>#V.E.</u>	FUNGAL STRAINS a	<u>WT</u>
g/ml	48	Lulwerthia sp. SC 73	0.
ç/ml	119	Leptosphaeria maritima RZ 312	1.
;/ml	210	Myrothecium sp. SC 87	1
ç/ml	'n	Trichoderma viride SC 51	2.
		<sup>a</sup> With culture collection accession	number
is inoculu	m for each	<sup>b</sup> Dry weight per unit volume; 2 ml experimental flask.	were us
the same o	ne ml pipette,	<sup>C</sup> Number of viable elements in one determined on cornneal agar sprea	drop fr dplates
		d Too many to count; inestimable.	

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Results.

ANOVA results of the four experiments are given in Tables 6-9 for nitrogen content (N%; arcsin transformation) and crude protein increment ( $\Delta$ CP) of the fermentation products, and treatment means are graphed in Figs. 2-7. In each case, the treatment means are graphed only if the ANOVA indicated that the treatments had

Table 10.

from 30 Two general patterns emerge from Figs. 2-7: (1) decrease in salinit  $_{3}$  > low  $0/\infty$  to 5  $0/\infty$  brings about increase in N% and  $\Delta$  CP, and (2) high NH<sub>4</sub>N N% and  $NH_4NO_3$  > low  $(NH_4)_2$  SO<sub>4</sub> with respect to their influence on increasing 4A, Fig.  $\Delta$  CP. These patterns, however, are clear in only a few instances (e.g. Fig. th other 3, 8% bran). Bran concentration was involved in significant interactions of In most treatments in every case but one (Fig. 7); no generalizations are possible. eralized cases, the interacting effects of two or all three treatments prevent generalized (e.g. Figs. 2,3, and 6).

y in 8% <u>Leptosphaeria maritima</u> RZ 312 converted wheat bran most efficien i and 3). bran, at 5 o/oo salinity and with the high concentration of  $NH_4NO_3$  (Figs. iable 3), The  $\triangle$  CP% (40) was near the highest of the bran culture screen results culture and the  $\triangle$  CP, mg/ml, (5.4) was higher than that of the highest of the bran s in the screen results (4.3). This was in contrast to the performance of this funce lower first bran culture screen (N% = 4.1,  $\triangle$  CP % = 14.5; Table 2) at concentration of  $NH_4NO_3$  (0.24%) and  $KH_2PO_4$  (0.006%) of that experiment.

ent overL. maritimawas the only one of the four fungi tested to show improvecultureits culture screen performance.Lulworthiasp. SC 73 (Fig. 4) matched ii 5 o/ooscreen performances (Tables 2 and 3), but it achieved this only in mediaation ofsalinity, as opposed to the 15 o/oo media of the culture screen.Concen $NH_4N0_3$  $NH_4N0_3$  had no effect on N% or  $\triangle CP$ , but  $(NH_4)_2 S0_4$  gave lower N% that

See Figures Mean values with 95% confidence intervals of nitrogen percentage and change in determine how the bran fermentations might be optimized with regard to protein production. Results for <u>Leptosphaeria</u> <u>maritime</u> RZ 312, for which all experimental treatments (bran concentration, salinity, and nitrogen source) and all crude protein content of fermented wheat bran from an experiment designed to interactions between and among them produced significant ( $ar{P}$ , Type I error < 0.05) effects on both nitrogen percentage and crude protein change. 2 and 3. Table 6.

				ka,t					
	2% B <sup>C</sup>			5% B			8% B		
	s o/oo <sup>b</sup>			<u>S o/oo</u>			<u>S o/co</u>		
'n	15	30	ŝ	15	30	S	15	30	
 3.87	3.73	4.01	4.62	4.35	4.22	4.92	4.21	4.26	
4.05	3.66	4.04	4.12	4.04	4.06	4.28	4.26	4.06	
3.73	3.60	4.20	4.24	4.17	4.09	4.12	3.71	3.96	
			ACP	e,f					
2% B <sup>C</sup>				5% B				8% B	
 00/0			ומ	0/00				<u>S 0/00</u>	
 15	30	Ś		15	30		Υ	15	30
 1(0%)	3.0(4%)	65.0(31	1%) 4	3.0(20%)	32.0(15%	)   136	.0(40%)	41.5(12%)	6.5(2%)
 5(5%)	0.1(0%)	31.0(15	5%) 5	3.0(25%)	10.0(5%)		.0(18%)	34.5(10%)	17.0(5%)
0(7%)	16.0(19%)	43.5(20	0%) 3	0.5(14%)	17.0(8%)	50,	5(15%)	6.5(2%)	11.5(3%)

LONN

LONS

NNIH

Table 6 (continued).

- a Nitrogen percentage
- b Salinity
- c Bran concentration
- <sup>d</sup> HINN = high NH<sub>4</sub>NO<sub>3</sub> concentration; LONN = low NH<sub>4</sub>NO<sub>3</sub>; LONS = low (NH<sub>4</sub>)<sub>2</sub> SO<sub>4</sub>.
- Change in crude protein in mgs., with percent change in parentheses.
- f Confidence interval not computed; these means are for two
  replicates only.



(S o/oo) on nitrogen content (N%) of fermented wheat bran. See Table 6 for further details. Leptosphaeria maritima R2312. Means which were significantly affected by treatments (P, Type I error < 0.05). The interacting effects of bran concentration, nitrogen source, and salinity Figure 2.



Table 7. Mean values with 95% confidence intervals of nitrogen percentage and change in crude protein content of fermented wheat bran from an experiment designed to determine how the bran fermentations might be optimized with regard to protein production. Results for <u>Lulworthia</u> sp. SC 73, for which bran concentration, salinity, nitrogen source, and the interaction between nitrogen source and bran concentration produced significant (P, Type I error < 0.05) effects on nitrogen percentage, and bran concentration, salinity, and the interaction of bran concentration and salinity produced significant effects on crude protein change. See Figure 4.

∆CP<sup>e</sup>

					<u>N %</u>	a. 	
					HINNd	LONN	LONS
1.	5	4.60 <u>+</u> 0.13		2	4.68 <u>+</u> 0.31	$4.67 \pm 0.41$	4.20 + 0.40
5 o/co <sup>D</sup>	15	4.27 <u>+</u> 0.18	вĩ <sup>с</sup>	5	4.19 <u>+</u> 0.64	4.22 <u>+</u> 0.33	4.17 ± 0.38
	30	3.90 <u>+</u> 0.15		8	4.08 <u>+</u> 0.33	4.04 <u>+</u> 0.30	4.06 <u>+</u> 0.40

			<u>S_0/00</u>	
		5 ·	15	30
	2	11.0 <u>+</u> 2.6 (13%)	4.3 <u>+</u> 9.9 (5%)	3.0 <u>+</u> 4.8 (4%)
Β%	5 8	33.0 <u>+</u> 13.9 (15%) 30.2 <u>+</u> 16.0 (9%)	16.8 <u>+</u> 13.5 (8%) -9.7 <u>+</u> 13.4 (3%)	-8.3 <u>+</u> 6.7 (4%) -34.5 <u>+</u> 13.9 (10%)

a Nitrogen percentage

b Salinity

c Bran concentration

<sup>d</sup> HINN = high NH<sub>4</sub>NO<sub>3</sub> concentration; LONN = low NH<sub>4</sub>NO<sub>3</sub>; LONS = low  $(NH_4)_2$ SO<sub>4</sub>.

<sup>e</sup> Change in crude protein in mgs., with percentage change in parentheses.



 Figure 4. Lulworthia sp. SC73. Means which were significantly affected by treatments (P, Type 1 error < 0.05). A. The effect of salinity (S o/oo) on nitrogen content (N%) of fermented wheat bran. B. The interacting effects of bran concentration (Bran %) and nitrogen source on nitrogen content. C. The interacting effects of salinity and bran concentration on crude protein increment (ACP, mgs). See Table 7 for confidence intervals and %A CP.

Table 8. Mean values with 95% confidence intervals of nitrogen percentage and change in crude protein content of fermented wheat bran from an experiment designed to determine how the bran fermentations might be optimized with regard to protein production. Results for <u>Myrothecium</u> sp. SC 87, for which concentration of bran, salinity, and the interaction of bran concentration and nitrogen source produced significant (P, Type I error < 0.05) effects on nitrogen percentage, and bran concentration and the interaction between bran concentration and salinity produced significant effects on crude protein change. See Figure 5.

N % a

ρ

					HINN <sup>d</sup>	LONN	LONS
	5	4.41 <u>+</u> 0.13		2	4.71 <u>+</u> 0.24	4.59 <u>+</u> 0.16	4.33 <u>+</u> 0.11
S o/oo <sup>b</sup>	15	4.23 <u>+</u> 0.13	B% <sup>℃</sup>	5	4.37 <u>+</u> 0.28	4.25 <u>+</u> 0.21	4.37 <u>+</u> 0.18
	30	$4.40 \pm 0.13$		8	4.11 <u>+</u> 0.25	$4.11 \pm 0.30$	4.29 <u>+</u> 0.10

			<u>ACP -</u>	
			<u>S 0/00</u>	
		5	15	30
	2	16.5 <u>+</u> 8.4 (19%)	17.3 <u>+</u> 8.0 (21%)	12.7 <u>+</u> 8.1 (15%)
в %	5	-7.5 <u>+</u> 23.0 (4%)	1.8 <u>+</u> 8.8 (1%)	9.8 <u>+</u> 7.6 (5%)
	8	0.3 <u>+</u> 20.5 (0%)	-27.2 <u>+</u> 22.0 (8%)	-15.3 <u>+</u> 11.1 (5%)

a Nitrogen percentage

b Salinity

c Bran concentration

- <sup>d</sup> HINN = high NH<sub>4</sub>NO<sub>3</sub> concentration; LONN = low NH<sub>4</sub>NO<sub>3</sub>; LONS = low (NH<sub>4</sub>)<sub>2</sub> SO<sub>4</sub>.
- e Change in crude protein in mgs., with percentage change in parentheses.



<u>Myrothécium</u> sp. 5C8/, MeahS Which were significantly affected
 by treatments (P, Type I error < 0.05). A.: The effect of salinity</li>
 (S o/oo) on nitrogen contenti(N%) of fermented wheat bran. B.
 The interacting effects of bran concentration (Bran %) and nitrogen source on nitrogen content. C. The interacting effects of salinity and bran concentration on crude protein increment (ACP, mgs).
 See Table 8 for confidence intervals and % ACP.

Table 9. Mean values with 95% confidence intervals of nitrogen percentage and change in crude protein content of fermented wheat bran from an experiment designed to determine how the bran fermentations might be optimized with regard to protein production. Results for <u>Trichoderma viride</u> SC 51, for which bran concentration, salinity, the interaction between bran concentration, salinity, and nitrogen source produced significant (P, Type I error < 0.05) effects on nitrogen percentage, and bran concentration and nitrogen source produced significant effects on crude protein change. See Figures 6 and 7.

		2% в <sup>с</sup>			5% B			8% B	
		<u>S o/oo<sup>b</sup></u>			<u>s o/oo</u>			<u>s o/oo</u>	
	5	15	30	5	15	30	5	15	30
HINN	3.94	3.59	3.72	3.95	3.82	4.00	4.07	4.04	3.96
LONN	3.79	3.94	3.95	3,90	3.57	3,72	3.87	3.73	3,99
LONS	3.56	3.54	4.04	4.17	3.84	3.89	4.18	4.02	3.84

∆CP<sup>e</sup>

	2	-17.9 <u>+</u> 3.2 (21%)	HINN	-32.8 <u>+</u> 7.7 (15%)
в %	5	-37.0 <u>+</u> 5.7 (17%)	LONN	-40.1 <u>+</u> 11.2 (19%)
	8	-49.3 <u>+</u> 9.3 (14%)	LONS	-31.3 <u>+</u> 8.0 (15%)

a Nitrogen percentage

b Salinity

c Bran concentration

- <sup>d</sup> HINN = high NH<sub>4</sub>NO<sub>3</sub> concentration; LONN = low NH<sub>4</sub>NO<sub>3</sub>; LONS = low (NH<sub>4</sub>)<sub>2</sub> SO<sub>4</sub>.
- e Change in crude protein in mgs., with percentage change in parentheses.

f Confidence interval not computed; these means are for two replicates only.



Trichoderma viride SC51. Means which were significantly affected by treatments ( $P_{1}$ , Type I error < 0.05). The interacting effects of bran concentration, nitrogen source, and salinity (S  $o/\infty$ ) on nitrogen content (N%) of fermented product. See Table 9 for further details. Figure 6.



Figure 7. <u>Trichoderma viride</u> SC51. Means which were significantly affected by treatments (P, Type I error < 0.05). The effects of bran concentration (Bran%) and nitrogen source on loss of crude protein (ACP, mgs). See Table 9 for confidence intervals and %ACP.

at the 2% level of bran concentration, where N% produced with  $NH_4NO_3$  was highest. At the optimal salinity level of 5 o/oo, 2% bran gave lower  $\triangle$  CP than 5 and 8% bran.

<u>Myrothecium</u> sp. SC 87 which had exhibited the highest N% and  $\triangle$  CP of the culture screens (Tables 2 and 3), gave much lower N% and  $\triangle$  CP in its 3 x 3 x 3 ANOVA experiment. Its highest significantly affected mean N% was 4.7 at 2% bran and the higher NH<sub>4</sub>NO<sub>3</sub> concentration. As with <u>Lulworthia</u> sp., (NH<sub>4</sub>)<sub>2</sub> SO<sub>4</sub> gave a lower N% than NH<sub>4</sub>NO<sub>3</sub> at this bran concentration. The 5 o/oo and 30 o/oo salinities gave higher N% than 15 o/oo. Highest  $\triangle$  CP, 17.3 mg (21%), was given by 2% bran, and there appeared to be little effect of salinity at this bran concentration though it had marked effects at the 5 and 8% bran levels. Nitrogen source had no effect on  $\triangle$  CP.

As in the case of <u>Myrothecium</u> sp., <u>Trichoderma viride</u> SC 51 (Fig. 6) exhibited much lower N% (maximum mean of 4.2%) in the 3 x 3 x 3 ANOVA experiment than it had in the second culture screen (5.3%; Table 3), though its response in the first culture screen (4.0%; Table 2) was lower than its ANOVA experiment maximum. In most cases, the higher  $NH_4NO_3$  concentration gave higher final N%, though this situation was reversed at 15 - 30 o/oo salinity with 2% bran and not apparent at 5 o/oo - 5% and 30 o/oo - 8%. At all three bran concentrations, highest N% was given by  $(NH_4)_2 SO_4$ ; this was at 30 o/oo salinity with 2% bran, and at 5 o/oo with 5 and 8% <u>bran</u>. The A CR reduces was all negative *JEig.* 70 is oncentration of NH; NOV Pro-

than either the high  $NH_{\mu}NO_3$  or  $(NH_{\mu})_2 SO_{\mu}$ .

following.
 Visual observations made during harvest of the flasks included to clearing of
 <u>The (NH, ), S0, flasks of peither SC 73, por BZ 312 exhibited the mark increase</u> in the induid phase evident in the NH NO. Hasks. There was a marked the induid phase evident in the NH NO. Hasks. There was a marked the shes from
 conidium formation by SC 51 as the percentage of bran decreased (the wincreased, the 2% bran fermentation were bright green). As the percentage of bran

the size of the mycelial pellets formed by SC 87 decreased markedly.

Since neither Myrothecium sp. nor Trichoderma viride performed in a fashion corresponding to its bran culture screen results, it is possible that these discrepancies were due to the standardized harvest time (7 days) of the  $3 \times 3 \times 3$  ANOVA experiment. Both of these fungi are fast-growing (Table 5). As can be seen from Table 4, maximum & CP% for Trichoderma viride from Microferm runs was +25%, at a bran concentration, nitrogen source and level, and salinity which gave negative  $\Delta$  CP (>10%) in the ANOVA experiment. When the set of conditions indicated as optimal for N% and  $\Delta$  CP by the ANOVA experiment were tried in a Microferm run with T, viride, results quite comparable to the ANOVA results were obtained (N% = 4.1 and  $\triangle CP = -31\%$ ) (Tables 4 and 9 and Figs. 6 and 7). This suggests that the ANOVA experiment showed best results for that combination of treatment levels which caused the fungus to grow less quickly into the later stationary phase (when conidium production--conidia are not efficiently harvested by 42 µ m filtration--and other loss of cell nitrogen takes place). Comparison of runs 35 and 36 (Table 4) with Table 8 shows another distinct lack of correspondence, but in this case, the cause is not clearly a time-of-harvest problem as with T. viride; Microferm runs 22, 25, and 27 (Table 4) for Myrothecium sp. did not correspond with the culture screen results either (Table 3), so that scale-up difficulties of some sort appear to have been involved with this fungus. Likely candidates as sources of these difficulties are differences in autoclaving of materials and in availability of dissolved oxygen during fermentation.

The control figures for N% and  $\triangle CP$  (Table 10) appear to show that the products from flasks with  $NH_4NO_3$  without fungi lost more CP and were lower in N% than those with  $(NH_4)_2 SO_4$ , no inorganic nitrogen, and no inorganic nitrogen + <u>Lulworthia</u> sp. However, the 5 means for neither N% nor  $\triangle CP$  are significantly different from one another (1-way ANOVA). It is interesting to note that when no

Table 10. Control values for the 3 x 3 x 3 ANOVA experiments designed to determine how the wheat bran fermentations might be optimized with regard to protein production.

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	HINND	LONN	LONS	NON
% N <sup>C</sup>	2,91	3.04	3.16	3.15
∆CP <sup>d</sup>	-106.0 (-31%)	-93.0 (-27%)	-77.5(-23%)	-77.5 (-23%)

8 %,	15	0/00,	Non,	Lulworthia	sp.	<u>SC 73</u>
% N				3.	.10	
∆CP				-(	66.5	(-20%)

2 % Sucrose, 15 0/00, 0.45 % NH<sub>4</sub>NO<sub>3</sub>

	<u>RZ 312<sup>e</sup></u>	<u>SC 73</u>	<u>SC 51</u>	<u>SC 87</u>
N %	3.12	3.10	4.31	7.38

- <sup>a</sup> Flasks treated like all others in the experiment, except that no fungal inoculum was added; medium of 8% wheat bran and 15 o/oo salinity; other components as in all other flasks.
- <sup>b</sup> HINN = 1.38% NH<sub>4</sub>NO<sub>3</sub>; LONN = 0.69% NH<sub>4</sub>NO<sub>3</sub>; LONS = 0.54% (NH<sub>4</sub>) $_2$ SO<sub>4</sub> Non = no added nitrogen-

 $^{\rm C}$  %N = nitrogen percentage in washed and dried product.

- d Change in crude protein from original (mg.) with percent change in parentheses.
- <sup>e</sup> Culture collection accession numbers for fungi used; see text for binomials.

inorganic nitrogen was added, no increase in N% or prevention of loss of CP took place with <u>Lulworthia</u> sp. (though the fungus obviously grew). This may indicate that the protein which is solubilized during autoclaving of the wheat bran is largely unavailable, at least to <u>Lulworthia</u> sp. It is evident from comparison of N% for mycelium controls (in which  $C/NH_4 - N =$  that of the bran media with the high concentration of  $NH_4NO_3$ ) with the N% of corresponding bran flasks, that the mycelium of some of the fermented bran products must have had higher N% than the mycelial controls. This is especially true in the case of <u>Leptosphaeria maritima</u> RZ 312 (unfermented wheat bran, N% = 2.67; mycelial control, RZ 312 N% = 3.12; in 8% bran, bigh NH<sub>4</sub> NO<sub>200</sub> 5 ploo salinity, RZ 312 N% = 4.92).

9. Feeding tests, 1974

Materials and methods.

In June, 1974, we began a series of four feeding experiments with penaeid shrimp (pink - <u>Penaeus duorarum</u>, and white - <u>Penaeus setiferus</u>). The same concrete tanks as used by Caillouet <u>et al</u>. (1975b) were used (Section 5). The layout of the Turkey Point shrimp mariculture facility where these experiments were conducted is given by Tabb <u>et al</u>. (1969) and updated by Caillouet <u>et al</u>. (1973). In all four of our 1974 artificial detrital feeding experiments (series F74), water was obtained from adjoining South Biscayne Bay and filtered through 5  $\mu$  m filter-cloth bags into the concrete tanks. The tanks were not prefertilized and the water was not aged. Flowing water systems were not used. Overnight aeration was conducted only when dissolved oxygen concentration approached 3 ppm (which it did only twice, during F74A). All shrimp used in the experiments were hatched and raised through their larval stages at the Turkey Point nursery according to the methodology described by Tabb <u>et al</u>. (1972). In each experiment fungal-fermented wheat bran products were tested as feed, with controls of unfermented bran (all experiments), commercial flaked feed (F74A, F74D), fertilization with inorganic fertilizer (F74C, F74D), and no

feed at all (F74A, F74B). The feeds were always thoroughly wetted with tapwater before they were added to feeding tanks, so that they would immediately sink and become available to the shrimp. All fermented feeds were prepared in the Microferm fermentor and processed as described above (Section 7). In most cases, fermentation products were dried at 55°C and ground either by mortar and pestle or in a Wiley mill. When this was not the case, products were harvested and washed as usual, but then refrigerated rather than dried.

Results.

The results of the F74 Experiments are given in Tables II and I2 and Figs. 8-II. Average final fresh weights (AFFW) were analyzed by 1-way ANOVA followed by Studekn-inEvenation Exists SINK) description and anong means, with confidence level = 95%. Logarithmic transformation was necessary for F74A and F74B, in which variance was inhomogeneous (Bartlett's test, p = 0.05). AFFW was approximately equal to average fresh weight gain in experiments F74A, B, and D due to the very small average initial weights of these experiments. Fresh weight gains were used in the statistical analysis of F74C, since the average initial fresh weight in that experiment was much larger than in the others.

One of the clearest conclusions to be derived from these results is that there were differences from tank to tank not due to intended treatments. These often caused marked, statistically significant differences in shrimp yields and AFFW. For this reason, means are not calculated for combined results of replicate tanks. Rather, replicate results are shown separately, so that the range of mean results is perceivable. Two major possible reasons for the unintended differences among tanks are differences in nutrient retention among tanks following previous experiments, and differences in the meiofauna and microbial flora (autotrophic and heterotrophic) which developed along with the shrimp, due to inoculum differences and nutrient retention differences. Yield values in Tables II and I2 are fresh weights, and feed

Table 11.	Results of shrimp-feeding experiments in the F 74 Series:
	F 74A and F 74B, with pink shrimp (Penaeus duorarum).
	Treatments are identified in Table 13. See Figures 8 and 9.

Duration, days Salinity range, o/oo Temperature range, <sup>O</sup> C Stocking density/m <sup>2</sup> Ave. initial wt. g	¥         58         32-38         26-31         15         0.010	<u>Ave. final wr.</u>	<u>Yteld g/m<sup>2</sup></u>	Survival %	Feed efficiency	60 35-38 27-31 15 0.005	Ave. final wt.	Yield g/m <sup>2</sup>	Survival %	Feed efficiency
Treatment	1	0.29	4.1	93		1	C.35	3.6	70	
	1	0.17	2.2	83	<del></del>	1	0.21	2.2	70	
	2	1.19	16.1	90	9	2	0.94	10.8	77	14
	2	0.92	12.9	93	11	2	1.27	14.6	77	10
	3	0,72	10.8	100	13					
	3	0.48	6.2	87	23					
	4	1.35	20.2	10 <b>0</b>	7					
	4	1.27	15.2	80	10					
	5	0.54	7.2	90	20					
	5	0.40	5.6	93	26					
	6	0.76	11.3	100	13					
	6	0.60	8.7	97	17					
	7	0.46	6.9	100	21					
	7	0.47	7.0	100	21					
	8	0.29	2.5	57	30					
	8	0.29	2.9	67	50					
	9	0.31	4.6	97	32	10	0.62	7.5	80	20
	9	0,43	6.4	100	23	10	1.02	11.7	77	13



Figure 8. Average final fresh weights, with 95% confidence intervals, of pink shrimp from feeding experiment F74A. Feeding treatments: I-no feed; 2-unfermented wheat bran; 3-autoclaved wheat bran; 4-Purina shrimp chow; 5-wheat bran fermented by <u>Trichoderma</u> <u>viride</u> SC51 (dried); 6-treatment #5, 90%, #4, 10%; 7-10% level, treatment #4; 8-treatment #5 (undried, cold-stored); 9-wheat bran fermented by SC87 (dried). Bars at base of graph join those means which could not be shown to be significantly different from one another (P, Type I error >0.05). Details in text and Table II.



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Duration, days	09 F 74C	<u>Ave. wt. gain. g</u> .	Gain, g/m <sup>2</sup>	Survival. %	Feed efficiency	<u>140</u>	Ave. final wt.	Yield, g/m <sup>2</sup>	Survival %	<u>Feed efficiency</u>
Salinity range o/oo	35-41					33-45				
Temperature range, <sup>o</sup> C	21-26					15-24				
Stocking density/m <sup>2</sup>	5					7.5				
Average initial wt, g	3.27					0.017				
Treatment	11	0.63	2.5	80		11	0,73	2.9	53	
	11	0.54	2.7	10 <b>0</b>		11	0,97	1.5	20	
	2	2.59	9.1	70	8	2	0.81	5.3	87	16
	2	1.85	9.2	100	8	2	Lost		<b>-</b>	
	12	1.26	6.3	100	12	4	1.23	8.0	87	11
	12	1.03	5.2	100	14	4	1.11	6.6	80	13
	13	2.07	9.3	90	8	15	0.79	5.1	87	17
	13	1.39	6.3	90	12	15	0.84	4.2	67	20
	14	0.74	2.9	80	26	16	0.68	4.1	80	24
	14	0.45	2.3	100	33	16	0.57	4.0	93	24
						17	0.79	3.5	60	28
						17	0.67	4.0	80	24



Figure 10. Average fresh weight gains, with 95% confidence intervals, of white shrimp from feeding experiment F74C. Feeding treatments: Il-regular fertilization; 2-unfermented wheat bran; 12-wheat bran fermented by <u>Pestalotia</u> sp. SC38 (dried); 13-treatment #3, 50%, treatment #2, 50%; 14-wheat bran fermented by <u>Curvularia</u> sp. SC86 (dried). Bars at base of graph join those means which could not be shown to be significantly different from one another (P, Type I error > 0.05). Details in text and Table 12.



Figure II.	Average final fresh weights, with 95% confidence intervals, of
-	pink shrimp from feeding experiment F74D. Feeding treatments:
	2-unfermented wheat bran; 4-Purina shrimp chow; 15-wheat bran
	fermented by Myrothecium sp. SC87 (undried, cold-stored), 50%,
	treatment 3, 50%, 16-wheat bran fermented by Trichoderma viride

- Table 13. List of treatments used in shrimp-feeding experiments of the F74 series. Compare with Figures 8-11 and Tables 11 and 12. Except where otherwise indicated, feeding levels were equal on a dry weight basis within experiments.
- 1. No feeding or fertilization.
- 2. Unfermented wheat bran. F74A, F74B: not ground; F74C, F74D: ground (Wiley mill) through 20-mesh screen. Mean CP% = 16.7.
- 3. Wheat bran, autoclaved, washed, and dried as in fermented wheat bran treatments, but <u>not</u> inoculated or fermented; ground in mortar and pestle. Mean CP% = 18.9.
- 4. Purina 'shrimp chow' ('Experimental Marine Ration 20'); fed as thick flakes as received. Ingredients include fish meal, soybean meal, ground wheat, brewer's yeast, dried whey, soybean oil, dicalcium phosphate, iodized salt, vitamins, minerals. CP% = 21.8.
- 5. Wheat bran fermented by <u>Trichoderma</u> viride SC 51. Dried at  $55^{\circ}$ C and ground by mortar and pestle. See text for CP% and  $\Delta$ CP%.
- 6. Treatment #5, 90%, and treatment #4, 10%.
- 7. Treatment #4, at 10% of its feeding level.
- 8. Treatment #5, but refrigerated rather than dried.
- 9. Wheat bran fermented by <u>Myrothecium</u> sp. SC 87. Dried at 55°C and ground by mortar and pestle.
- 10. Treatment #9, but refrigerated rather than dried. See text for CP% and  $\Delta CP$ %.
- 11. Fertilization by daily addition of a commercial fertilizer (18-24-16) such that daily addition of nitrogen was equivalent to that of the most nitrogenous of the other feeds in an experiment; pre-dissolved.
- 12. Wheat bran fermented by <u>Pestalotia</u> sp. SC 38; dried at  $55^{\circ}$ C, ground (Wiley mill) through 20-mesh screen. See text for CP% and  $\Delta$ CP%.
- 13. Treatment #12, 50%, and treatment #2, 50%.
- 14. Wheat bran fermented by <u>Curvularia</u> sp. SC 86; dried at 55°C, ground (Wiley mill) through 20-mesh screen. See text for CP% and ΔCP%.

Table 13 (continued)

- 15. Treatment #10, 50%, and treatment #2, 50%.
- 16. Treatment #5 (but ground (Wiley mill) through 20-mesh screen), 57%, treatment #2, 43%. Fed 13% more/day than other treatments in F74D. See text for CP% and ΔCP% of fermented portion.
- 17. Like treatment #16, but the fermented portion ground (Wiley mill) through a 60-mesh screen . Fed 13% more/day than other treatments in F74D. See text for CP% and ΔCP% of fermented portion.

efficiency = total dry weight of feed + total fresh weight of animals. The treatments used in the four feeding experiments are identified in Table 13.

Experiment F74A was begun in June and harvested in August (Table 11). Wheat bran fermented by <u>Trichoderma viride</u> SC 51 and by <u>Myrothecium</u> sp. SC 87 was tested as feed for postlarval pink shrimp. The fermented materials came from Microferm runs 19-22 and 24-26. As Table 4 shows, these materials varied in N% from 28.3 to 31.8 and in  $\triangle$  CP% from +5 to +25 for SC 51, and in  $\triangle$  CP% from +5 to +11 for SC 87 with N% = 31.6. The SC 51 material was fed both dried (mortar and pestle ground) and refrigerated. Other feeding treatments were unfermented, unground wheat bran, autoclaved, washed, dried and ground wheat bran, commercial flaked feed ("shrimp chow," Purina Experimental Marine Ration 20), a combination treatment of 90% SC 51 fermented bran and 10% commercial flakes, a commercial flakes treatment at 10% of the feeding level of the above flake treatment, and a no-feed control. The experimental feeds were fed at 5 gms per day on a dry weight basis.

The commercial flakes performed best in AFFW in both replicates (Fig. 8), although the higher of the unfermented wheat bran replicates was not significantly different from those of the commercial pellet replicates (lower than the highest one by a factor of only 0.9). The autoclaved and washed bran replicates were both lower in AFFW and total yield than the unfermented bran replicates, although one of the replicates of each fell in a group of three means which were not significantly different from one another. Thus, autoclaving of wheat bran does not, in itself, impart higher nutritive capacity to bran.

The higher AFFW of the replicates of the dried <u>T. viride</u> fermented-bran feed was only slightly higher than the AFFW of the higher replicate of the <u>Myrothecium</u> sp. product (significant difference was not detected between them), and both AFFW's were lower than the higher of the unfermented bran AFFW's by a factor of about 0.5. Both replicates of the refrigerated T. viride feed were as low in AFFW as the higher of the no-feed replicates, lower by a factor of 3 than the unfermented bran replicates. In total yield, the refrigerated <u>T</u>. <u>viride</u> feeds were lower than the higher of the no-feed controls, due to the markedly lower survivals, indicating a possible toxicity of the refrigerated <u>T</u>. <u>viride</u> material. It must be noted here that refrigeration problems apparently occurred during storage of this feed material, for bacterial spoilage was quite noticeable by the time of harvest for F74A. The addition of 10% of commercial flakes to the dried <u>T</u>. <u>viride</u> feed produced marked improvement in AFFW and total yield; the higher of the AFFW's of the mixedmaterial replicates was significantly different from and 1.4 times greater than that of the higher of the unmixed replicates, although it was still smaller than that of the higher of the undermented bran replicates by a factor of 0.6. That this improvement was not due to the commercial flake fraction alone, is shown by the fact that the AFFW of the higher of the replicates of the diet which consisted only of commercial flakes (at 10% of the daily feeding level of the other diets) was significantly lower than the AFFW of the higher of the mixed-diet replicates (by a factor of 0.6).

In July, experiment F74B was initiated. Experimental conditions were the same as those of F74A (Table II), with one important exception: the average initial fresh weight (5 mg) of the postlarval pink shrimp used was one-half that of F74A. Since other experimental conditions were little different between the two experiments, it was probably this factor that led to the reduced survival observed in F74B. Three feeding treatments were applied: wheat bran fermented by <u>Myrothecium</u> sp. SC 87 (product washed and refrigerated); unfermented, unground wheat bran; and a no-feed control. Feeding levels were the same as those of F74A (5 gm/day on a dry weight basis), and experiment duration (60 days) was approximately the same as that of F74A (58days). The fungal material came from Microferm runs 27 and 28 (Table 4), the products of which were very similar (in N% and  $\triangle$  CP%) to those of the Microferm runs (22 and 25) which yielded the <u>Myrothecium</u> - feed treatment of F74A.

The means of all treatment replicates were significantly different from one another in F74B. The AFFW's and total yields of the unfermented bran replicates of F74B were similar to those of F74A (Fig. 9), and the same was true of the no-feed controls; in both cases, total yields were slightly lower in F74B. The most marked difference between F74A and F74B was the performance of the <u>Myrothecium</u> feed. The AFFW of the higher of the two replicates of the refrigerated feed (F74B) was higher by a factor of 2.4 than that of the higher replicate of the dried feed (F74A). It was also lower than the higher AFFW of the unfermented bran in F74B by a factor of only 0.8. Similar figures applied to comparisons among the same treatments with regard to total yield. These results represented a marked improvement over the dried feed with respect to comparison on a higher-replicate basis to the unfermented bran treatments; the dried-feed AFFW of F74A was smaller by a factor of 0.45 than the AFFW of unfermented bran.

Experiment F74C was begun in August (Table 12). Salinities ranged higher and temperatures lower than in F74A and F74B, but the major differences in experimental conditions were that a different species of shrimp was used (white shrimp, <u>Penaeus setiferus</u>) and average initial fresh weight was much larger (value with 95% confidence interval =  $3.27 \pm 0.21$  gm). For this reason, average fresh weight gain (AFWG) and total fresh weight gain (TFWG) were compared among treatments. Also, stocking density was lower on the basis of number of individuals per m<sup>2</sup> (5), but much higher on the basis of fresh weight per m<sup>2</sup> (8.2gm, as compared to 0.1gm for F74A). Feeding level was 2.5 gm per day on a dry weight basis (approximately 15% of body weight at the beginning of the experiment).

feeding treatments. Two fungal-fermented feeds were tested, one produced by <u>Pestalotia</u> sp. SC 38 (Microferm runs 31 and 32) and <u>Curvularia</u> sp. SC 86 (Microferm run 29). In both cases, N% and  $\triangle$  CP% were lower than had been the case with the previously tested <u>Trichoderma</u> and <u>Myrothecium</u> feeds (Table 4). Both fungal feeds of F74C were dried and ground through a 20-mesh Wiley mill screen. One combination treatment wassured, an <u>SO SOM inviture of Pestalotia</u> stp. SC 38 officied material and unfermented bran.

The highest AFWG was that of the higher replicate for unfermented wheat bran (Fig. 10), but it was closely followed by that of the higher of the mixed-treatment replicates (lower by a factor of only 0.8, and not significantly different). The TFWG of the higher of the mixed treatment replicates was higher than that of the higher of the bran replicates. The higher replicate of the unmixed <u>Pestalotia</u>-fermented material had an AFWG which was smaller by a factor of 0.6 (significantly different) than that of the higher of the mixed-treatment replicates. The <u>Curvularia</u> replicates fell in a group of means not significantly different from one another, including the inorganic nitrogen controls, and the higher replicate of the <u>Curvularia</u> feed had an AFWG which was only 1.2 times higher than the higher inorganic nitrogen control. In TFWG, the <u>Curvularia</u> results were virtually the same as those of the inorganic nitrogen controls (Table 12).

Pink shrimp postlarvae were again used in experiment F74D, begun in September (Table 12). The average initial fresh weight (17 mg) was larger than that of F74B, but percent survival was again lower than in F74A, probably due to the combined effects of higher salinity range and lower temperature range. Stocking density was one-half that of F74A and F74B, and feeding level was similarly reduced (2.6 gm/day on a dry weight basis).

As in F74C, an inorganic nitrogen control and a 20-mesh unfermented wheat bran treatment were applied. The commercial flake treatment used in F74A was also

applied. The three fungal-fermented bran treatments were all mixtures with 20-mesh unfermented bran. <u>Myrothecium</u> sp. SC 87 material (refrigerated, not dried) was mixed 50-50% with unfermented bran. <u>Trichoderma viride</u> SC 51 material was dried and ground through two mesh sizes, 20 and 60. Each of these two sizes served as a separate feeding treatment, mixed at 57% fungal material to 43% unfermented bran. The mixed <u>Trichoderma</u> feeds were fed at 3 gm/day. The fungal materials were produced in Microferm runs 26 and 34-36 (Table 4). Runs 34-36 were lower in N% and markedly lower in  $\triangle$  CP% than previous runs with the same fungi. In addition, the refrigerated <u>Myrothecium</u> material was accidentally frozen prior to the beginning of F74D, and by the end of the experiment, it had suffered noticeable spoilage.

Because survival was markedly different among treatments and even among replicates, and because there was an apparent marked effect of survival upon AFFW (Table 12), only those replicates with greater than 80% survival were compared by ANOVA and SNK (Fig. II). As in F74A, the commercial flake replicates were both higher in AFFW than those of the unfermented bran treatment; in F74D, the higher flake replicate was 1.5 times larger in AFFW and total yield than the one bran replicate (and significantly different from it). The unfermented bran AFFW and total yield, however, were virtually identical to those of the one <u>Myrothecium</u> + bran replicate with survival greater than 80%. The higher <u>Trichoderma</u> + bran treatment replicates with survival greater than 80% gave lower AFFW (though significant difference was not detected) and poorer feed efficiencies (by a factor of 1.4) than the higher-survival <u>Myrothecium</u> + bran replicate. There was no apparent difference between the two <u>Trichoderma</u> + bran treatments with different particle sizes.

10. Culture screening on delignified bagasse and straw

## Background.

Plant waste materials of minimal acquisition cost are, of course, the eventual target of our feed-development program. The agricultural wastes of least value are
those which do not readily serve as productive feeds for terrestrial farm animals, i.e., those which are high in lignified cellulose (Pigden and Heany, 1969). The lignin portion of these materials acts on a molecular level as a physical and/or chemical barrier to cellulose digestion, and thus devalues the materials as feed for ruminants. Bagaske is a good example of an agricultural waste material of this type. It is the remains of the sugar cane stalk after crushing to remove sucrose, and on a dry weight Jassis; constitute of JWS, callulose - themicallulose, 20%. Lignin, and 2% silice. (Scinivasan and Han, 1969). Materials with lignin and silica contents as high as these serve largely as inert fillers or roughage in feeding of ruminants (Van Soest, 1969). Wheat straw is also one of the lowest quality forages; like bagasse, it is very low in nitrogen content (0.4%), and addition of urea in itself has no effect on its digestibility for ruminants (Pigden and Heany, 1969).

Wheat straw and bagasse, then, are two typical lignocellulosic, low-value agricultural by-products. In addition, both are easily obtainable in South Florida, and bagasse is readily available in the warm tropical countries where shrimp mariculture development is taking place (Gross, 1973; Webber, 1975). Therefore, these two materials were chosen as substrates for a fungal culture screen.

The results of our earlier culture screen on bagasse (Section 6) had demonstrated the marked resistance to fungal decomposition of this material. Other workers had shown a similar response using cellulolytic fungi and bacteria (Chahal <u>et al.</u>, 1969; Srinivasan and Han, 1969). As stated above, wheat straw is degraded only to a small extent by rumen microorganisms. However, in the case of both materials, physical and chemical treatment of the materials which alters the lignin-cellulose physical/chemical relationship can greatly improve their degradability (Callihan and Dunlap, 1971; Toyama and Ogawa, 1972; Wilson and Pigden, 1964). Therefore, we applied a delignification (this word is used here to mean alteration of the lignin-cellulose relationship, rather than complete removal of lignin) treatment to the bagasse and straw which was to be used in the culture screen.

Materials and methods.

The treatment was a "dry" method and involved a combination of conditions used in several other delignification studies (Ghose and King, 1963; Han and Srinivasan, 1968; Pew and Weyna, 1962; Toyama and Ogawa, 1972; Wilson and Pigden, 1964). The oven-dried (55<sup>°</sup>C) materials were ground through a 20-mesh Wiley mill

and autoclaved (120°C) for one bour after thorough minimum at a 25% (whither we

n is closer to a wet solid than ned with tapwater over a 245 became clear (12 liters were then resuspended in 800 mls th tapwater until constant pH about 10 liters in each case; both bagasse and straw, the ginal dry weight. This was in rough the coarse filter cloth t 55°C and ground through a

availability, it was deemed
m the earlier bagasse screen.
j by the agar-plate growth
67), Trinci (1969, 1971), and
gal biomass measurement by
(1971). Their methods were
to degrade and convert the

(w/v) NaOH solution in deionized water. (This combine a liquid suspension.) When cool, the materials were  $v_{\mu}$  m hylon filter cloth until the liquid passing the filrequired for each of the materials). The materials w of 5% acetic acid, stirred for 60 seconds, and washed of the resuspended materials was reached (this requifinal pH: straw, 5.4; bagasse, 5.2). In the case delign-fication and washing resulted in a 62% loss of part due to the loss of finely divided solid materia during washing. The materials were then oven-drie 60-mesh Wiley mill screen.

suspension of the lignocellulosic materials (200.gms

Due to constraints of time, funds, and equipal desirable to use a different culture-screening method It was decided to try an experimental plan insp measurement methodology developed by Pirt (1966 Morrison and Righelato (1974), and the methods of direct observation of Warnock (1971) and Nagel-deB adapted to yield indices of degree and rate of abi

bagasse and straw substrates.

The fungi chosen for screening were selected for one or more of the following reasons: high frequency of occurrence on bagasse and/or straw before or during the shrimp-feeding experiment described in Section 5; description in the scientific literature (Sections 3 and 4) as highly cellulolytic and/or of high potential or indicated nutritive value; high nitrogen content produced in wheat bran fermentation or relatively high nitrogen content produced in bagasse fermentation during earlier culture screening (Section 6); isolation from estuarine or maritime cellulosic substrates and availability in our culture collection. Many of the fungi chosen were not taken all the way through the culture-screening procedure. In some cases, this was due to poor growth on the inoculum or test media, but in others, time constraints were the cause.

The experimental procedure was as follows. All cultures to be tested were inoculated onto fresh plates of commeal agar (CMA) made up with 15 o/oo seawater. These and all following plates were sealed with masking tape and incubated in darkness at  $25^{\circ}$ C; as soon as ample vigorous growth was produced on these CMA plates by each fungus, transfer was effected onto plates of 1/4-strength CMA with additional agar to bring it to 2% (1/4 CMA +A). These 1/4 CMA +A plates were of uniform agar media depth (20 ml/plate, 10 cm diameter plates) and were prepared at the same time and using the same batch of 15 o/oo seawater as were all the following plates. The transfer onto 1/4 CMA +A and all following transfers were done uniformly as follows: a sterile #1 cork borer (5 mm diameter) was used to remove a cylinder of agar from the plate to be inoculated; a cylinder was then taken from 5 mm behind the growing edge of the inoculum plate, inverted, and placed in the hole on the plate to be inoculated; a cylinder was intended.to.deplate.inocula...

of excessive endogenous reserves. The fungi were permitted to grow on these until they had reached the edges of the plates or until growth slowed. They were then

inoculated onto plates of agar media with incorporated delignified material. These

	<u>ຼຼາງຄທັງ ແພກສາແທດງານໄປປະຕິດສຳສະເພງກິດ ເດຍີດສ່ຽງເວລະ ແລະປະເຊ</u> າເດີເອີຍ. ການຂະຍາດ
ate of	KH2P04, 0.01% yeast extract, 2.0% agar, in 15 o/oo seawater. One
epared	delignified straw medium (DSA) and one of delignified bagasse (DBA) wa
ates or	for each test fungus. Again, these were allowed to grow to the edges of th
் நஜர்	
	grew very poorly or not at all and were eliminated from further testing.

ed toFrom these inoculum plates of delignified media, the fungi were transfDBA.the culture screen plates. Six media were used, two of which were DSA atDSACorresponding plates of media with non-delignified materials were preparedTotfeau...courumgua, aux sumstitic NBA)....Twn-Distort of screen plates.Six media were used, two of which were DSA atTotfeau...courumgua, aux sumstitic NBA)....Twn-Distort of screen plates.CMA (only Difco commeal agar added to the 15 o/oowater), and one of plain agarCMA (only Difco commeal agar added to the 15 o/oolia except bagasse or straw.medium (PA) having all the ingredients of the test...Three cylinders from the inoculum plates were juxta

multimum biologous and an environmentation process to be derived from the culture screen results.

Each test plate was read once daily after inoculation. Colony radial growth extent and rate (mm/hr) were read for three radii on each plate. Each of the radii bisected a sector which contained one of the inoculum cylinders. Plates were read until growth of 2 of the 3 sectors reached the edge of the plate, or until growth rate of the 3 sectors slowed by a total of 0.1 mm/hr over a 24-hour period. When this occurred, the plates were first stained with trypan blue in lactophenol, then flooded with a killing and fixing agent (FAA: ethyl alcohol (95%) - 50%, glacial acetic acid - 5%, formaldehyde (40%) - 10%, water - 35% (Sass, 1958)).

The stained and fixed plates were used to make a determination of biomass of fungus produced. A cylinder of agar was removed from 5 mm behind the leading edge of each of the 3 sectors on each plate. A 2 mm thick slice was taken from the top of each cylinder and placed on a microscope slide. An eyepiece with photographic reticle (Wild) having a transecting line was used to measure hyphal density. The eyepiece line was positioned so that it was perpendicular to the majority of hyphae in the field, and all surface hyphae crossing the line were then counted and recorded as hyphae per mm. Cover slips were then placed on the slices and the diameters of 5 randomly selected hyphae from each of the three sectors were measured. The 15 hyphal diameters and 3 densities were averaged to yield values of each for each of the test plates for each fungus.

Using these two measurements, an indicator was calculated of the fungal biomass produced, simply by multiplying mean hyphal density by mean hyphal diameter for both control (PA) and test media, and subtracting the value for PA from that of each of the test media. This is given as biomass index (BI) in Table 14 for those few fungi for which analysis was completed, alongside an indicator of potential

utility in the fermentation process to be developed (UI. = Bl.x maximum growth rate). stationary phase is also given in Table 14, but not included in idices (although it might logically have been) for this was the ned of the growth parameters measured.

r which complete analyses were conducted (Table 14), the 2 iC 14 and 134) and Chaetomium sp. SC 6 gave the highest BI's 578 respectively), and Chaetomium sp. SC 6 gave by far the . Surprisingly, only 4 of the 15 fungi produced higher BI on these, marked increases were shown by Chaetomium sp. SC 6, SC 125, and Trichoderma viride SC 51. This is likely to have Growth extent at earl the calculation of the least accurately deter

Results.

Of the 15 fungi species of Drechslera on DSA (728, 658, and highest BI (629) on DS DSA than on NDSA. C Pycnoporus sanguineu

		fungi capable (alkali-trea biomass.	e of effici ted) wheat :	ently convert: straw and baga	ing "delign asse into f	ified" ungal	
FUNCI <sup>a</sup>	SUBSTRATE	MAX. Growth <u>Rate<sup>c</sup></u>	GROWTH EXTENT CLASS <sup>d</sup>	MEAN HYPHAL DENSITY <sup>e</sup>	MEAN HYPHAL DIAM.f	BIS	<u>uip</u>
	PA	0.17	1	9.0	2.1		
	CMA	0.34	. 3	58.0	3.4	197	67
SC 6	NDS	0.40	3	74.5	4.1	305	122
	DS	0.40	3	121.9	5.4	658	263
	NDB	0.34	2	36.6	2.7	<b>98</b>	33
	DĿ	6.41	3*	91.3	6.9	629	257
	PA	0.29	3	11.4	2.1		
	CMA	0.37	3*	58,3	2.6	128	47
SC 8	NDS	0.35	3*	42.6	2.8	95	33
	DS	0.33	3	18,6	1.5	4	1
	NDB	9 <b>.31</b>	3*	25,2	1.8	21	7
	DB	0.31	3*	12.6	2.0	1	0
	PA	0.31	3*	9.3	3.1		
	CMA	0.35	3	70.9	4.1	262	92
SC 14	NDS	0.31	3	305.1	4.2	1253	388
	DS	0.33	3	302.7	2.5	728	240
	NDB	0.37	3*	36.6	4.5	136	50
	DB	0.30	2	87.1	3.6	284	85
	PA	0.25	1	23.4	2.2		
	CMA	0.36	3	78.1	3.4	214	77
SC 20	NDS	0.37	2	45.6	3.3	99	37
	DS	0.28	3*	43.2	2.1	39	11
	NDB	0.34	2	22,8	3.4	26	9
	DB	0.30	3*	30.6	2.3	19	6

Table 14. Results of a culture screen designed to find strains of

Table 14. (continued)

FUNCIa	SUBSTRATE <sup>D</sup>	MAX. GROWTH <u>RATE<sup>C</sup></u>	GROWTH EXTENT CLASS <sup>d</sup>	MEAN HYPHAL DENSITY <sup>e</sup>	MEAN HYPHAL DIAM. <sup>f</sup>	BIg	<u>ui</u> h
	PA	0.61	3*	8.7	4.5		
	CMA	0.83	3*	74.5	5,2	348	289
SC 51	NDS	0.79	3*	58.9	4.9	249	197
	DS	0.80	3*	106.3	3.9	375	300
	NDB	0.71	3*	20.4	3.7	36	26
	DB	0.75	3*	82.3	4.3	315	236
	PA	0.34	2	5.7	3.7		
	CMA	0.32	3*	27.0	3.5	73	23
SC 77	NDS	0.34	1	96.7	4.2	<b>38</b> 5	131
	DS	0.36	1	53.1	4.3	207	75
	NDB	0.35	3*	35.4	4.5	138	48
	DB	0.30	3*	33.6	3.5	97	29
	PA	0.25	2	5.3	2.4		
	СМА	0.50	3*	18.6	4.8	<b>7</b> 7	38
SC 90	NDS	0.44	2	30.0	3.7	<b>9</b> 8	43
	DS	0.35	3*	20.4	2,5	38	13
	ND B	0.31	3	6.3	3.5	9	3
	DB	0.32	3*	13.5	3.3	32	10
	PA	0.37	3*	58.9	4.9		
	CMA	0.39	3*	120.1	4.2	216	84
SC 98	NDS	0.43	3*	77.5	5.1	107	46
	DS	0.41	3*	97.3	3.5	52	21
	NDB	0.37	3*	93.7	3.3	21	8
	DB	0.38	3*	141.2	3.2	163	62

# Table 14. (continued)

FUNGIª	SUBSTRATE <sup>b</sup>	MAX. GROWTH RATE <sup>C</sup>	GROWTH EXTENT <u>CLASS<sup>d</sup></u>	MEAN HYPHAL <u>DENSITY</u> <sup>e</sup>	MEAN HYPHAL DIAM, f	BIS	<u>ui</u> h
	PA	0.18	1	24.9	1,6		
	CMA	0.24	2	38.7	2.3	49	12
SC 100	NDS	0.30	2	91.9	2.1	153	46
	DS	Lost	. 1	15.6	1.5		
	NDB	0.24	3*	33.6	1.7	17	4
	DB	0.14	1	20.7	1.4	-11	
	PA	0.06	0	9.0	2.3		
	CMA	0,19	2	39.9	2.7	87	17
SC 101	NDS	0.45	3*	75.7	4.9	350	158
	DS	0.43	3	51.6	4.5	212	91
	NDB	0.41	3*	27.0	5.4	125	51
	DB	0.45	3*	54.7	3.9	193	87
	РА	0.19	3	17.4	2.1	<del>~ ~</del>	
	CMA	0.19	0	91.3	1.8	128	24
SC 114	NDS	0.25	3*	94.6	1.9	143	36
	DS	0.22	1	104.5	2.0	172	38
	NDB	0.21	3*	22.8	3.4	41	9
	DB	0.19	3	69.7	3.7	221	42
	PA	0.28	3*	9.9	5.7		
	CMA	0.13	2	60.7	4.0	186	24
SC 122	NDS	0.30	2	111.7	4.6	457	137
	DS	0.28	3*	48.6	3.8	128	36
	NDB	0.35	2	36.3	4.7	114	40
	DB	0.29	3*	28.8	3.5	44	13

FUNGIª	SUBSTRATE	MAX. GROWTH <u>RATE<sup>C</sup></u>	GROWTH EXTENT CLASS <sup>d</sup>	MEAN HYPHAL DENSITY <sup>e</sup>	MEAN HYPHAL DIAM.f	BIg	<u>ui</u> h
	РА	0.22	2	70.2	3,2		
	CMA	0.26	2	303.9	2.8	626	163
SC 125	NDS	0.25	3*	193.4	2.5	259	65
	DS	0.25	. 3*	217.4	2.9	406	101
	NDB	0.21	2	91.0	3.5	94	20
	DB	0.24	3*	164.6	2.5	187	45
	PA	0.54	3*	25.5	2.1		
	CMA	0.40	3*	69.7	2.6	128	51
SC 133	NDS	0.35	3*	94.9	2.2	155	54
	DS	0.38	3*	80.5	2.0	107	41
	ND B	0.36	3*	34.8	1.9	13	5
	DB	0.38	3*	54.7	1.3	18	7
	PA	0.20	2	9.9	2.6		
	CMA	0.20	1	73.0	4.5	303	61
SC 134	NDS	0,31	2	158.6	6.3	973	302
	DS	0.30	2	133.3	4.6	587	176
	NDB	0.26	3*	7.8	2.3	-8	
	DB	0.25	2	45.1	3.7	141	35

<sup>a</sup> Fungal strains listed by accession (culture) number; SC 6--Chaetomium sp.; SC 8--Scopulariopsis(?) sp.; SC 14--Drechslera sp.; SC 20--Zythia sp.; SC 51--Trichoderma viride; SC 77--Dendryphiella salina; SC 90--Aspergillus niger; SC 98--Pestalotia sp.; SC 100--Sporormia sp.; SC 101--Chaetomium sp.; SC 114--Aspergillus terreus; SC 122--Alternaria longissima; SC 125--Pycnoporus sanguineus; SC 133--Schizophyllum alneus; SC 134--Drechslera sp.

## Table 14. (continued)

- <sup>b</sup> PA = plain agar (control); CMA = cornmeal agar (Difco); NDS = non-delignified straw; DS = delignified straw; NDB = non-delignified bagasse; DB = delignified bagasse. All media 2.0% agar (Difco) (except CMA), 0.25% NH<sub>4</sub>NO<sub>3</sub>, 0.09% KH<sub>2</sub>PO<sub>4</sub>, 0.01% yeast extract (Difco), 2.9% test substrate (except CMA), in 15 o/oo seawater.
- <sup>c</sup> Maximum growth (diametric extension) rate observed over a 24-hour period, in mm/hr.
- d 0 = 0-10 mm; 1 = 10-20 mm; 2 = 20-30 mm; 3 = 30-40 mm; 3\* = rate of growth not declining when growing edge had reached plates edge. Plates were flooded with fixative (FAA) when growth rate declined by an average of 0.03 mm/hr or more over a 24-hour period.
- <sup>e</sup> Across a 1 mm-long transect made perpendicular to the growing hyphae 5 mm from the growing edge.
- f In µm, average of fifteen randomly-selected hyphae.
- 8 BI = biomass index = (mean hyphal diameter (MHDI) on test substrate x mean hyphal density (MHDE) on test substrate) - (MHDI on plain agar x MHDE on plain agar).
- h UI = apparent utility index = BI x maximum growth rate.

been an effect of excessive alkali treatment and thorough washing having removed soluble nutrients, including otherwise available hemicellulose (Wilson and Pigden, 1964; Garcia-Martinez et al., 1974) which may have positive effects on cellulolysis and growth (Basu and Ghose, 1960; Nilsson, 1974).

Of the 11 fungi which produced higher BI on NDSA than on DSA, the highest values and most marked differences were shown by the <u>Drechslera</u> spp. (SC 14 and 134) and <u>Alternaria longissima</u> SC 122. To the contrary, II of the 15 fungi produced higher BI on DBA than on NDBA, highest values and most marked differences being shown by <u>Chaetomium</u> sp. SC 6, <u>Trichoderma viride</u> SC 51, and <u>Aspergillus terreus</u> SC 114. Of the four which did not, only <u>Alternaria longissima</u> SC 122 produced a markedly higher BI on NDBA than on DBA, and the other three had low BI's on both media. Thirteen of the 15 fungi had higher BI values on the straw agars than on the corresponding bagasse agars. Of the two which were otherwise, <u>Pestalotia</u> sp. SC 98 had highest BI on DBA.

Maximal radial growth rates were much less evidently affected by the test substrates than hyphal diameter and especially hyphal density, so that within a fungal test group, UI values reflect BI values. However, because of its very rapid growth rate (approximately 0.75 mm/hr, 24-hr mean, on all media except PA), <u>Trichoderma viride</u> SC 5I produced the highest UI on DSA (300), followed by <u>Chaetomium</u> sp. SC 6: 263, and <u>Drechslera</u> sp. SC 14: 240, and was second only to <u>Chaetomium</u> sp. SC 6 on DBA (SC 6: 257, SC 51: 236; next highest was 87 for <u>Chaetomium</u> sp. SC 101). Highest UI on NDSA was 388 (<u>Drechslera</u> sp. SC 14) followed by <u>Drechslera</u> sp. SC 134 at 302, and the next highest was <u>Chaetomium</u> sp. SC 101 at 158.

The values of UI and BI for CMA relative to these values for the other test substrates varied among the fungi tested. Some fungi exhibited highest values on CMA (e.g. Pycnoporus sanguineus SC 125 and Pestalotia sp. SC 98), while others exhibited values on CMA as low or lower than those on NDB, which was, in general, the poorest substrate (e.g. <u>Dendryphiella salina</u> SC 77 and <u>Chaetomium</u> sp. SC 101).

### DISCUSSION

### II. Fungal fermentation of wheat bran

Wheat bran was not chosen as a substrate for our initial fungal fermentations because ot its potential for scaled-up commercial production of feed. Its cost is substantially higher (5 times or more) than that of the nearly entirely lignocellulosic agricultural by-products since its relatively high soluble carbohydrate (53%) and protein (17%) content give it value as a feed fraction and bakery product. Rather, we chose it because, of the four plant materials tried as artificial detrital materials by Caillouet <u>et al.</u> (1975b; Section 5), it was the only one which : (1) supported marked growth of shrimp when fed without treatment (Caillouet <u>et al.</u>, 1973, 1975a, b); and, <u>memory importentials (2) negotited marked growth of function</u>.

We chose wheat bran in order to determine on a preliminary basis: (1) whether the concept of feeding of fungal fermented materials was a practicable one; and (2 whether the yield of shrimp producible with untreated wheat bran could be improved by fermentation, beyond the size satisfactory for a commercial bait-fishery concern.

Our culture screen results indicate the extent to which the protein content of wheat bran can be changed by fungal fermentation (under the specific condition imposed) (Tables 2 and 3). Conditions of the two screens differed in several respect (Section 6), but perhaps most importantly in that phosphate concentration an especially inorganic nitrogen concentration were markedly higher in the secon screen. Probably as a consequence, the percentage of nitrogen in the product wa pushed higher in the second screen (Fig. 1); the mutability of nitrogen content of fungal mycelium as influenced by the C/N ratio of the growth medium is well know (e.g. Cowling, 1970; Lilly, 1965; Section 3). It is probably fair to suggest that the higher nitrogen contents reflect increases in true protein content, for Levi and Cowling (1969) and Graham (1971) found that decrease in the C/N ratio of growth medium leads to increase in non-nucleic nitrogen of mycelium, but does not result in - mancherease oidermetric view lages costing of amountation

There was a lesser effect on the percentage change in crude protein; the greatest fraction on mose rungi restert reunderendet to crude protein by 0.2.49% in both, screens, though the second screen produced fewer high losses and more high gains. The difference in extent to which the nitrogen content of the mycelium of a given fungus is mutable is shown by the three strains of fungi which were tested in both screens; one (Lulworthia sp. SC 73) produced nearly identical results in both screens, while the other two (Trichoderma viride SC 51 and Pestalotia sp. SC 38) showed increases in percentage of nitrogen of 8.3 and 12.5 respectively. These increases may also have been partially due to increased phosphate concentration. The  $P0_4$  concentration used in the first culture screen (0.006%) is likely to have been limiting (Child <u>et al.</u>, 1973), unless  $P0_4$  present in the wheat bran could alleviate the shortage.

Comparison of our wheat bran culture screen results with the results of other investigators who have attempted protein production by fermentation of substrates of high starch content is difficult for the following reasons. (I) In some cases, results are incompletely presented, due to the proprietary nature of the research. (2) Entirely soluble media were used, rather than solubles + particulate suspension--thus separation of mycelium from unconverted material was possible. (3) Determination of mycelium production by nitrogen analysis of the fermented wheat bran products wassnot possintesible describe the original protein content of wheat bran is considerable.

(4) Differences of opinion exist among investigators with regard to the best means of determining protein content of fungal mycelia (e.g. Christias <u>et al.</u>, 1975; Coleman, 1972; Herbert <u>et al.</u>, 1971; Solomons, 1973) resulting in the use of several methods (Lowry, Biuret,  $\propto$  -amino nitrogen, total amino acids, total nitrogen); conversion of

figures derived by one method to those derived by another are impossible. (5) Differing methods of presentation of results have been used (biomass/carbohydrate utilized, protein/carbohydrate utilized, protein/volume of fermentation liquid, etc.). (6) Although wheat bran is a common growth medium used in the production of fungal enzymes (Codner, 1969; Ghandi et al., 1974; Underhofler et al., 1947; Wang et al., 1974) we know of only one very recent report of its use for the production of fungal biomass (Von Hofsten, 1974). With regard to item (4), it is perhaps best in our particular case to use total nitrogen x 6.25, since the major part of the error in calculating true protein from total nitrogen lies in the contribution of chitin nitrogen. At least a portion of the chitin may be nutritively valuable to shrimp (Section 4).

Results of several studies of protein production by fungal fermentation of largely-starch or simple-sugar substrates are given in Table 15. When substrates such as these are utilized for production of fungal biomass, and only the resultant

# mycelium is collected the crude wate in content of the products can capse avite bigh If, however, the solid undigested remainder of the

since	included in the fermentation product, the upper end of the range is lower (38
yn of	non-nitrogenous, unfermented plant structural materials (some combin
f our	cellulose, hemicellulose, lignin, pectin, residual starch) remain. The resul
s, are	wheat bran fermentations, the products of which include undigested bran fi
other	much the same in terms of range of crude protein content as those us
Table	substrates in particulate suspension. The values for protein production give
n and	15 are dependent upon concentration $(w/v)$ of substrate used in the ferment
em to -	upon ability of the fermenting microorganism(s) to efficiently convert
olume	microbial biomass at those concentrations. Thus, they are related to
a/unit	production efficiency (biomass produced/unit volume of fermentation m
	time), though the figures in Table 15 do not include growth-rate information.
other	The wheat bran values (and probably the value for most of 1

The wheat bran values (and probably the value for most of

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yloid	bran	
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б	8 U 8 8	n uo
Results	simple-	mentati
51		
Table		

			Crude Pro	otein % <sup>C</sup>	Prote	ein Yield <sup>d</sup>
Reference	<u>Substrate</u> b	#Strains	Low	High	Low	<u>High</u>
iolomons. 1973	nr <sup>e</sup>	"hundreds"	NR	74	NR	NR
alanghe et al 1964	soybean whey, L,9	4	33	64	0.5	4.6(+1.3)
raham. 1971	glucose, L, NA <sup>f</sup>	ຕາ	35	62	0.5	2.9
Christias et al., 1975	glucose, L, NA	4	39	58	NR	NR
<u></u> 3ednarski et al., 1971	milk whey, L,O	4	16	57	3.9	23.3 <sup>1</sup>
dtchfield, 1968 <sup>h</sup>	several, NA	33	e,	54	en En	NR
Reade et al., 1974	cassava, L, 0.4	"extensive screening"	NR	48	NR	NR .
Brook et al., 1969	cassava, S <sup>g</sup> , 0.4	28	16	42	1.0	2.8
<u> </u>	molasses, L, 4.6	1	25	38	2.5	8.7(+5.9)
Gray, et al., 1964	glucose, L, NA	175	13	31	1.2	3.4
Gray & Abou-El-Seoud, 1966a	sweet potato, S,7	Q	15	38	1.0	5.9(+0.9)
Culture screen II	wheat bran, S,17	25	21	34	6.7	12.8(+4.3)
Chahal & Cheema, 1971, 1972	rice, S,7	20	21	33	0.4	2.5(+1.5)
Alcrofermj	wheat bran, S,17	4	25	32	2.5	10.6(+2.1)
BFIEk	wheat brau, S,17	4	22	31	2.7	19.0 (+5.4)
Reade et al., 1972	barley, S,Il	23	œ	31	NR	2.8 (+0.6)
<u> </u>	rice, S,7	14	24	30	1.8	4.6 (+3.0)
Grav & Abou-El-Seoud, 1966c	sugar beets, S,3.4	5	22	29	1.2	4.2 (+3.1)
Culture screen i	wheat bran, S,17	27	15	57	1 1	9.9 (+0.6)
Gray & Abou-El-Seoud, 1966b	cassava, S,2.8	5	13	20	1.2	3.0 (+2.2)

Table 15 (continued)

- entirely liquid media are listed first, followed by those of liquid suspensions Listed in decreasing order of maximal crude protein content; fermentations of of particles. æ
- Listed with an indication of medium type (L = entirely liquid, S = suspension of particles), and the initial crude protein percent. م,
- c Of final dried product.
- In mg/ml of medium. Numbers in parentheses indicate change in mg/ml from original. σ
- e Not reported or determinable from the reference cited.
- f Not applicable.
- Particulates were removed after fermentation and not included in the protein analysis. 50
- h A review paper.
- 1 It is not clear whether the mycelial product was washed.
- J Results of 10.5-liter fermentations; see Table 4.
- Results of the bran fermentation improvement experiment; see Tables 11-12 and Figures 8-11. ч.

particulate-suspension fermentations in Table 15) are likely to include protein remaining or converted from original substrate protein in addition to that produced from inorganic nitrogen. Therefore, production of protein over quantities previously present is given in parentheses. Since some of the original protein is undoubtedly lost from the solid phase during the fermentations (Section 8), and this was not measured, the actual protein production values are not known. The highest values for fungal protein production over that originally present in wheat bran are as high as the highest of the other values of protein production given in Table 15, except for those of Bednarski <u>et al.</u> (1971). Higher values not yet reported may well have been achieved in the recent extensive research projects of Reade <u>et al.</u> (1974), Solomons (1975), and Worgan (report due for publication in J. Sci. Fd. Agric.).

The protein production figures of Bednarski <u>et al.</u> in Table 15 are an order of magnitude higher than most of the others given. Prefermentation of milk whey by bacteria, followed by mold fermentation, is reported to have been the reason for these high values, but confirmation of these results is needed: the concentration of easily fermentable sugars and conversion efficiencies necessary to have achieved these results would have to have been quite high; the production results of El-Akher <u>et al.</u> (1974), working with yeast fermentations of milk whey, do not approach those of Bednarski <u>et al.</u>, washing of the fermentation product is not reported by Bednarski <u>et al.</u>

# 12. Effect of simultaneous modification of fermentation conditions

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1.1.1.

protein content and amino acid balance of fungal biomass. Pinto's work clearly demonstrated that use of  $(NH_{4/2}) \leq 0$  could have a provider influence on content of the sulphy fryl amino acids (one of the most common deficiencies of fungal protein--Chiao and Peterson (1953); Litchfield (1968); Rhodes <u>et al.</u> (1961); Thatcher (1954) ). Other workers (Hueck and Hazeu, 1969; Meyers 1968; Sguros <u>et al.</u>, 1973) had concluded that  $NH_4NO_3$  could serve as well as or better than several other nitrogen sources in providing for good growth of fungity. Therefore, we examined the effects of zero doublesses.

of NH NO<sub>3</sub> concentration and one level of  $(NH_{4})_2 SO_4$  in the fermentation medium on the final nitrogen percentage and crude protein increment produced in fungalfermented wheat bran. Gray <u>et al.</u> (1963) and Jones and Irvine (1971) had shown that salinity of the fermentation medium could have marked effects on mycelial and protein yield, so we examined this phenomenon. Finally, we examined the effects of concentration of substrate, which is one of the most direct determinants of volume production efficiency (Section II), and thereby one of the most important factors in economic evaluation of any developmental fermentation process (Callihan and Dunlap, 1971; Gray, 1971).

The results of our comparison of nitrogen sources with regard to their effectiveness in stimulating protein production (Section 8) reflect the findings of other investigators who have examined the question of optimal nitrogen source for fungal fermentations (Table 16). Interaction is common among the effects of type of nitrogen source and several other variables (type of carbon source, concentration of carbon source, medium pH and medium buffering, other medium constituents, and, especially, species of fungus used). Also, optimal nitrogen source may be different depending upon the goal of the developers of the fermentation process (i.e. whether optimized nitrogen percentage of final product, optimal protein production, or optimal biomass production is desired). It is interesting to note, in the case of two fungal strains in the present study (Lulworthia sp. SC 73 and Myrothecium sp. SC 87),

	•		
Reference	Production & Substrate <sup>a</sup>	Fungi	Order of Effectiveness
Sekeri-Pataryas <u>et al</u> ., 1973	🄏 CP-carob extract	Aspergillus niger	$(\mathrm{NH}_4)_2 \mathrm{SU}_4 > (\mathrm{NH}_2)_2 \mathrm{CO}$
	DW "	H	$(NH_4)_5 SO_4 = (NH_2)_2 CO_4$
Chahal & Cheema, 1972	% CP & DW-rice	Pericillium crustosum	$KNO_3 > (NH_2) CO > NH_2 CL> NH_2 NO_3$
Shukla & Dutta, 1967	DW & CP/vol-molasses	khizopus sp.	$MH_{cL} > (DH_{c})_{2} SO_{4} > (MH_{2})_{2} CO_{6}$
Graham, 1971	DW & CP/vol-glucose	Rhizopus oligosporus	$(\text{NH}^4)^2 \text{sc}_4 \text{NH}_4 \text{cL} + (\text{NH}^2)^2 \text{cO} + (\text{NH}^4)^2 \text{HP}_6$
	DW-potato starch, pN 4	=	$(\text{NH}_{4})_{2}^{2}\text{HPO}_{4}^{4} > (\text{NH}_{4})_{2}^{2}\text{SO}_{4}^{2}$
	DW-potato starch, pH 6	-	$(\rm NH_4)^2 SO_4 > (\rm NH_4)^2 HPO_4$
	CP/vol-potato starch, $p\mathbb{R}$ 4	Ξ	$(NH_4)_{HPO_4}^{A} + (NH_4)_{SO_4}^{A} = 0.00$
	CP/vol-galactose	-	(NH <sub>4</sub> ) <sub>2</sub> S0 <sub>4</sub> >(NH <sub>4</sub> ) <sub>2</sub> HP0 <sub>4</sub>
Pinto, 1963	CP/vol-glucose	Heterocephalum aurantiacum	$(\text{NH}_2)_2^{\text{CO}} (\text{NH}_4)_2^{\text{SO}_4} > \text{NH}_4^{\text{NO}_3} > \text{NH}_4^{\text{CL}}$
	÷	Cladosporium sp.	$NH_{c}CL > (NH_{2})_{2}CO > NH_{a}NO_{3} = (NH_{2})_{2}SO_{4}$
	-	Spicaria elegans	$NH_{A}CL = (NH_{A})_{5}SO_{A} > (NH_{5})_{5}CC > NH_{A}NO_{3}$
	=	Linderina pennispora	$(\text{KH}_4)_2 \text{ so}_4 > (\text{KH}_2)_2 \text{ co}_{\text{MH}_2} \text{ cL}_{\text{MH}_4} \text{ NO}_3$
Yusef & Allam, 1967	DW-glucose	Chaetomium sp.	$(NH_2)_2 CO > N = NO_3 > (NH_4)_2 SO_4$
	=	Pestalotia gracilis	$NaNO_3 > (NH_2) 2CO> (NH_4) 2SC_4$
	Ξ	<u>Myrothecium</u> verrucaria	$NaNO_3 = (NH_2)_2 CO > (NH_4)_2 SO_4$
Verona <u>et al</u> ., 1973	% TAA-glucose	Aspergillus niger	$\overline{NH}_{2}CN > (\overline{NH}_{2})_{2}CO > \overline{NH}_{4}NO_{3} > \overline{NaNO_{3}}$
	=	Penicillium frequentans	$(\text{NH}_2)_2 \text{CO>NH}_4 \text{NO}_3 \text{>NH}_2 \text{CN>NaNO}_3$

The effectiveness of types of nitrogen source in stimulating fungal production processes. Table 16.

Reference	Production & Substrate <sup>a</sup>	Fung1	Order of Effectiveness
Aguihotri, 1964	DW-sucrose	Aspergillus quadrilineatus	$NaNO_3 > (NH_2)_2CC > (NH_4)_5C_4$
	DW-starch	=	(NH <sub>2</sub> ) <sub>2</sub> CO>NaNO <sub>3</sub> >(NH <sub>2</sub> ) <sub>2</sub> SO <sub>2</sub>
Sguros and Simms, 1963	DW-glucose	Halosphaeria mediosetigera	$KNO_{3} > KNO_{3} > (NH_{4})_{3}CO_{3} > NH_{4}NO_{3} > NH_{4}CL > (NH_{4})_{5}SO_{4}$
	", buffered, pH <sup>R.2</sup>	=	$(\mathrm{NH}^{\mathrm{V}})^{3}\mathrm{SO}_{\mathrm{m}}^{\mathrm{m}}\mathrm{SO}_{\mathrm{m}}^{\mathrm{m}}\mathrm{NH}^{\mathrm{m}}\mathrm{SO}_{\mathrm{m}}^{\mathrm{m}}^{\mathrm{m}}}\mathrm{SO}_{\mathrm{m}}^{\mathrm{m}}\mathrm{SO}_{\mathrm{m}}^{\mathrm{m}}^{\mathrm{m}}\mathrm{SO}_{\mathrm{m}}^{\mathrm{m}}^{\mathrm{m}}\mathrm{SO}_{\mathrm{m}}^{m$
Jones and Irvine, 1971	CP/vol-cellulose	Corollospora cristata	(NH <sub>2</sub> ), CO>KNO <sub>2</sub> >NH <sub>2</sub> NO <sub>2</sub> >NH <sub>2</sub> CL
	Ξ	<u>Dendryphiella</u> salina	$(NH_2)_5 CO^{-1} KNO_3^{-1} NH_2 NO_3^{-1} NH_4 CL$
	2	Cladosportum herbarum	(NH <sub>2</sub> ) , CO>NH, NO <sub>2</sub> > NH, CL>KNO <sub>3</sub>
	=	Lulworthfa sp.	$(NH_2)_5 CO > KNO_3 = NH_4 CL > NH_6 NO_3$
Hueck and Hazeu, 1969	radial growth	5 common cellulolytic species	NH <sub>A</sub> NO <sub>3</sub> =NaNO <sub>3</sub> > (NH <sub>2</sub> ) <sub>2</sub> CO>NH <sub>6</sub> CL
	cellulolytic activity	÷	NH_NO3> (NH,),CO>NH,CL>NaNO3
Chahal and Gray, 1968	CP/vol-wood pulp	44 species	(NH <sub>2</sub> ) <sub>2</sub> CO>KNO <sub>2</sub> >NH <sub>2</sub> CL
El-Kersh <u>et al</u> ., 1973	cellulase-CM cellulose	Phoma glomerata	$NH_{A} KO_{3} = NANO_{3} NH_{A} CL > (NH_{2})_{3} CO$
	=	<u>Rhizoctonia</u> solani	NaNO <sub>1</sub> >NH <sub>2</sub> NO <sub>1</sub> > (NH <sub>2</sub> ) <sub>2</sub> CO>NH <sub>2</sub> CL
Siu, 1951	cellulolysis	Aspergillus fumigatus	$(\mathrm{NH}_{\mathrm{A}})_{\mathrm{S}}\mathrm{SO}_{\mathrm{Z}}^{\mathrm{S}}\mathrm{Na}\mathrm{NO}_{\mathrm{T}}\mathrm{S}(\mathrm{NH}_{\mathrm{S}})_{\mathrm{S}}\mathrm{CO}_{\mathrm{T}}^{\mathrm{S}}$
	E	Humicola grises	$(\mathrm{NH}_{3})_{5}\mathrm{CO}^{3}(\mathrm{NH}_{4})_{5}^{3}\mathrm{SO}_{4}^{2}\mathrm{NaNO}_{3}$
	z	Aspergillus terreus	$NaNO_3 > (NH_4)_2 SO_4$
	•		
	a % CP = crude protein	percentage; DW = dry weight; CP	/vcl = crude proteín per volume
	of medium; ZTAA = per	centage of total amino acids.	

.

Table 16. (continued)

that although nitrogen source had a marked effect on final percentage of nitrogen in the fermented product, there was no detectable effect upon the absolute change in crude protein, meaning that the deficiencies in production of a highly nitrogenous product were balanced by difference in mycelial biomass produced or original material lost.

Clearly it is of little value to determine optimal nitrogen source for fungal fermentations without examining simultaneously the effects of other potentially interacting variables. However, two generalities which appear upon examination of the references cited in Table 16 are: (1) as Jones and Irvine (1971) point out, when urea,  $(NH_2)_2$  CO, has been tested, it has often performed as well as or better than other simple nitrogen sources; (2) as Nicholas' review (1965) and Sguros and Simms' (1963) work show, when ammonium salts are to be used, it is advisable to use buffered media. The present study did not include examination of applicability of these generalities to wheat bran fermentations; it is quite possible that higher levels of protein production might have been achieved had urea been used as a nitrogen source, or had better buffered media been used with the ammonium salts used.

Final percentages of nitrogen and crude protein yields of the present experiment appeared to depend more heavily on type of nitrogen source than on C/N ratio except in the case of <u>Leptosphaeria maritima</u> RZ 312 at the highest bran concentration, when the lower C/N gave markedly higher final nitrogen percentage and crude protein increment. That C/N had no effect on final nitrogen percentage of the <u>Lulworthia</u> products was to be expected, since this fungus had shown nearly equivalent results in the two culture screens, which had utilized media with much different C/N ratios (51 in culture screen I and 14 in culture screen II).

The two C/N ratios used in our experiment (14 and 28, considering only ammonium nitrogen--see Section 8; these values fall to about 7.5 and 10 when the bran nitrogen is included) lie in or near the range (5-25) given by Litchfield (1968) as

optimal for fungi in general with regard to fungal protein production. The C/N ratio must be interpreted with caution in media including complex substrates such as wheat bran, especially when ammonium and nitrate ions are present; not all of the organic carbon (Von Hofsten and Von Hofsten, 1974) or nitrogen of the wheat bran is available to fungi, and ammonium ion uptake depresses or prevents nitrate ior uptake (Nicholas, 1965), so that the ratio of available carbon to available nitrogen is not



It in both studies, the obligately marine fungi showed the otein production as salinity decreased from 30 o/oo to about 5 :ern was clearer than the one observed with type of nitrogen here too, salinity had significant interacting effects with the .es (in five of the eight possible cases, considering both final | crude protein increment).

were detected for the modified concentrations of wheat bran ght of eight), and significant interaction was detected in seven y general conclusion which can be reached, and this applies to int variables examined, is that in experimentation designed to inditions for a given developmental fermentation, one must ependencies exist among conditions which can be varied, and em together, as many at a time as is practicable, if one is to n production levels for given fungal strains. For example, if rotein production capacity of <u>Leptosphaeria maritima</u> RZ 312, source at 2% bran and 30 o/oo salinity, we would have  $S0_{\mu}$  produced optimal performance (Fig. 3). Then, varying

interesting to note clearest increases in o/oo. This general source (Section 8), 1 other treatment var nitrogen percentage Significant eff in every tested case of these cases. The all three of the trea determine optimum determine where int syste natically alter discover optimal pro we had examined th varying only nitrog concluded that (NH salinity at 2% bran with  $(NH_4)_2 SO_4$ , we would conclude that 30 o/oo permitted best performance. Finally, varying bran concentration at 30 o/oo and with  $(NH_4)_2 SO_4$ , we would conclude that bran concentration had little effect, and miss the peak performance at 8% bran with  $NH_4NO_3$  and 5 o/oo salinity!

If apparatus for continuous culture is available, the recently reported method of media optimization of Mateles and Battat (1974) might be adaptable to improvement of fungal fermentations.

## 13. Fielding of fermented wheat bran to penaeid shrimp



I¢/Ib. (Callihan and Dunlap, 1971); (2) wheat bran is not readily available in the Central and northern South American countries where commercial mariculture development is now centered (Webber, 1975); (3) wheat bran is composed of both easily formentable material (starch and protein) and highly refractile material (native cellulosics) (Caillouet et al., 1975; Von Hofsten and Von Hofsten, 1974). Caillouet et a.A <u>brvc a proximanetalinatysteinmbicshows a approximated ap</u> sources (Basu and Ghose, 1960; Bravery, 1968; Hueck and Hazeu, 1969; King and <u>Eggins, 1973; Zeltin, 1970; and especially Tashpulatov and Teslinova, 1974).</u>

Item (3) from the above paragraph has an important bearing on interpretation of the results of our shrimp-feeding tests. Because of the relatively high starch (wheat starch was shown by Forster and Gabbott (1971) to be very effectively assimilated by prawns) and protein content, wheat bran, fed directly to penaeid shrimp with no treatment whatever, will support the growth of shrimp to a size and at a rate satisfactory for a commercial bait fishery (Caillouet et al., 1973, 1975a, b). Performance of wheat bran, in terms of yield of shrimp produced, was often only slightly lower in both Caillouet's experiments and in the present experiments, than commercial (or developmental commercial) pelleted or flake feeds (Glencoe pellets, - Annias consisted soullase. Burias Experiments' Marise & Ration 20) . In Caillovet's experiments, yields with wheat bran ranged from 69 to 100% of yields with commercial feeds. Any fermentation of the bran will necessarily reduce the available energy per unit of original starting material (Von Hofsten, 1975), and since much of the dignified certholesc and hermized bleec will not be converted without pretreatment (several reports in Gould, 1969; Rogers and Spino, 1973), this fraction will be higher in the fermented product than in untreated bran. The cellulosic fraction of the bran is no more available as nutritive matter to penaeid shrimp than it is to the fungi. Cellulose has been used in shrimp feeds as an inert filler--e.g. Andrews et al. (1972); Forster and Gabbott (1971) found that calories of even a finely ground purified cellulose were assimilated by prawns to only a small extent. Therefore, on a unit dry weight basis, fermented bran will have a lower quantity of nutritively valuable material and available energy than untreated bran. Since the feeding treatments used in our feeding studies were applied at equivalent rates of dry weight per day, the animals receiving fermented bran were receiving less nutritive material than those receiving untreated bran.

Our 14-liter fermentations of wheat bran (Table 4) were conducted largely forproduction of material for feeding to shrimp, as was the case in the paper fermentation and chick-feeding study of Crawford et al. (1973); also in parallel with their work, the large-scale fermentations yielded lower quantities of protein per unit weight of original substrate in most cases, than had similar small (flask) scale fermentations, due to greater loss of dry weight with respect to the original. Loss of efficacy in yields of fermentations due to scaling up is common and to be expected in developmental studies of this kind (Gray and Abou-El-Seoud, 1966a; Hishinuma et al., 1972; Solomons, 1969). It is due to subtle differences in preparation and sterilization of media, in preparation of inoculum, and in oxygenation and other transport phenomena (Solomons, 1972). Systematic analysis of the variables involved in the 14liter fermentations was not conducted and more than one variable was often changed from run to run. Therefore, reasons for scale-up problems were not pinpointed. It is important to note that where crude protein produced was less than in small-scale fermentations, and more of the original weight was lost to respired gas on solubilization, the remaining product would contain a greater fraction of the cellulosic material not nutritively valuable to penaeid shrimp.

The results of our feeding experiments exhibit the lack of replicability among feeding treatments which is to be expected when experiments are conducted in outdoor facilities under semi-natural conditions. Under these conditions, interacting uncontrolled variables such as development of microbial and meiofaunal populations, weather effects, development of parasite and pathogen problems, etc., are not prevented from imposing marked and differing effects on shrimp growth. Problems of this sort are discernible in the results presented for other studies of this kind, for example, by Caillouet <u>et al.</u> (1973) and Parker and Holcomb (1974), and Zein-Eldin and Meyers (1974) encountered and discuss the problem. Until these natural variable

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factors can be better controlled, it is the higher levels of production achieved which should be noted, for these represent indications of levels attainable when some measure of control over experimental variables is accomplished; for this reason, we did not combine means for replications of treatments in our feeding experiments.

The range of responses to fungal-fermented feeds in our experiments ranged from very poor yields of shrimp due to a possible effect of toxicity, to slightly higher yields than unfermented bran. The yields using cold-stored <u>Trichoderma viride</u> material seemed to indicate toxicity, for they were lower than yields when no feed was supplied, due to heavier mortality. Cases of toxicity have been reported for <u>T</u>. <u>viride</u> (e.g. Mirocha and Christensen, 1974), but so have cases of good performance of <u>T. viride</u> mycelium as feed material (e.g. Church <u>et al.</u>, 1972). Improper refrigeration certainly could have been the factor responsible for the poor performance of this particular feed; when the <u>T. viride</u> material was dried, excessive mortalities did not occur.

The dried <u>T</u>. <u>viride</u> product gave poor shrimp yields relative to unfermented bran (feed efficiency  $\geq 20$ ) except when combined with commercial flake material (90% + 10%, respectively), but best feed efficiency was still only 0.7 that of unfermented bran. Two other dried fermented products, those of <u>Myrothecium</u> sp. and <u>Curvularia</u> sp., also gave feed efficiencies greater than 20. It is quite possible that the drying process used was at least in part responsible for the low nutritive capacity of these products. Bauer-Staeb and Bouvard (1973) found that heating and drying had very marked negative effects on the availability of yeast-cell proteins to digestive enzymes, and Spicer (1973) points out that excessive heat processing can reduce the availability of amino acids of fungal protein. Bressani (1968) and Kihlberg (1972) discuss the problem of processing of single-cell protein, they point out that treatments such as heating and drying can impair nutritive capacity by reduction of amino acid availability and/or loss of functional quality (e.g. vater absorption capacity). Barlocher and Kendrick (1973) found that detritivorous amphipods would not accept dried fungal mycelium, though fresh mycelium was avidly eaten.

After harvest of F74C, which utilized large and easily dissected shrimp, contents of the foregut and hindgut were examined. It was observed that the fibrous bran flakes were passing through the shrimp largely undigested, and that fungal material which had been dried onto the flakes was also passing through undigested (the percentage which went undigested was not determined). When the <u>Myrothecium</u> product was refrigerated rather than dried (and spoilage did not take place), a marked improvement in performance in terms of maximal shrimp yield was observed (Table II, treatments 9 and 10). Drying of the products is, of course, much more desirable than cold storage in terms of development of an economically desirable product, and further experimentation may demonstrate that other methods of drying, or drying of some other developmental fungal feeds, do not detract from nutritive capacity: Drying lass improved the nutritive value of single-cell protein in other cases (Kihlberg, 1972; Mogren <u>et al.</u>, 1973). See also Regnault <u>et al.</u> (1975) in this regard.

The refrigerated <u>Myrothecium</u> product (F74B) gave a maximal shrimp yield and average final fresh weight per shrimp which were only 0.8 those of the maximal yield of untreated bran of F74B. If we consider the question of relative quantities of nutritive value between the fermented and unfermented bran in this case, using conservative calculation of conversion of wheat bran to mycelial biomass, we find that unfermented bran contains 1.3 - 1.5 times as much nutritive material as fermented bran on a dry weight basis. In making these calculations, it is assumed that the portion of the wheat bran left unconverted by the fungus was also unavailable to the shrimp (see above, this section) and that the portion of the bran which was accessible to the fungus was converted at 40-50% efficiency. If these speculative calculations are fair approximations, then the fungal product may have actually given higher feed-conversion efficiency than the unfermented product, based on weight of nutritive matter added. Experiments involving feeding of purely fungal materials (no residual unfermented matter) or with more efficiently converted substrates (less residual unfermented matter) would be necessary to verify this conjecture.

In order to partially correct the differences in quantity of nutritive matter among fermented and unfermented bran feed in F74C and F74D, we applied feeding treatments of 50% fungal material and 50% unfermented bran. We did so for the further reason that plant starch has been found to be a valuable component of artificial penaeid diets (Andrews and Sick, 1973), and wheat bran starch and other nutritive components, as discussed above, obviously serve well in this capacity. The result of this mixing of fermented and unfermented feeds in F74C was to improve the maximal yield of shrimp, using a Pestalotia product, from 0.7 that of unfermented bran to slightly better than equivalency (Table 12). If half the smaller end of the correction-factor range (1.3) discussed in the above paragraph is applied to this yield. . Jeed. afticinogy-bacomer-casely-squal\_townbat\_bad\_beensenkingsdwrith. dae- Brniae Experimental Marine Ration 20 in F74A. Feed efficiency was slightly lower in F74C than in F74A for unfermented bran, indicating that the Pestalotia + unfermented bran feeding treatment performed on a par with the Purina product, on the basis of nutritive matter fed (assuming that the ratio of shrimp yield from unfermented bran to shrimp yield from the Purina product would have been the same in F74C as it was in F74A).

The results of F74D demonstrate the marked effect which temperature (and salinity?) can have upon shrimp growth (Zein-Eldin and Griffith, 1969): feed efficiency for wheat bran rose to twice its previous low value, and mortalities were, in general, higher than in F74A and F74C. Yield values for wheat bran and the Purina product were much lower than in previous experiments, and this was not entirely due

to the lower stocking density, for average final fresh weights were also distinctly lower. The mixtures of cold-stored <u>Myrothecium</u> product and dried <u>Trichoderma</u> product both performed less well than wheat bran in terms of shrimp yield, though the <u>Myrothecium</u> + unfermented bran product produced a yield which was only 4% less than the yield from unfermented bran. In addition to the ambient temperature problems, refrigeration problems occurred with the <u>Myrothecium</u> product in this experiment and probably further obscured real differences between this ard other feeding treatments. Under the conditions of this experiment, particle size in the range tested (20 - 60 mesh) had no evident effect on shrimp yield from dried Trichoderma material.

The shrimp yields of our F74 series of feeding tests are low and the feed efficiencies high when compared with some of the results of other workers who have investigated growth of penaeids on artificial diets (Table 17). They are, however, comparable to the findings of Caillouet <u>et al.</u> (1975b) developed in the same concrete tanks as used in the present study. That they are a bit lower than Caillouet's findings is probably due to the fact that Caillouet used inorganic fertilizer <u>in combination</u> with his wheat bran feeding treatments, and we did not do so. There are two basic reasons for the low yields: (I) among the penaeid shrimp, <u>Penaeus duorarum</u>, used in F74A, B, and D, is one of the least attractive species in terms of growth potential under the conditions imposed by a mariculture facility (Broom, 1973; Parker and Holcomb, 1974; compare with Forster and Beard, 1974); (2) the facilities and methodology used in outgrowth of shrimp at Turkey Point have consistently brought about low yields when compared with other experimental facilities (see Krantz and

\_Norris\_1976), Our results for these reasons and other experimental difficulties and other experimenta

inequivocally that fungal feeds can or should be ither, they suggest strongly that potential for ermented feed exists, especially in view of the

faults discussed above, do not state used in mariculture of shrimp. I development of a successful fungal

investigators.	are included.	-D.
shring as reparted by several	eding artificial preparations	e listed for experiments F 74A
wrowth of peansid	Only results of fe	Present results are
Table 17.		

						GAD d	Ш рај	υĺ
Reference	01	ipecies <sup>a</sup>	1GP b	Inc	<b>F</b> 0	(MA-MI) IH	ĪH	3
Broom, 1969	ы В В	<u>tiferus</u>	49-80	0.1 hect. ponds	10.0	0.97 (2.70-19.78)	4.9	3.0
		Ξ	31-48	20 m <sup>2</sup> ponds	0.21	0.41 (1.46-2.42)	14.8	5.4
	<u>P</u> .	tecus	60-100	0.1 hect. ponds	0.11	0.70 (0.54-11.97)	9.7	1.7
		5	60	20 m <sup>2</sup> ponds	0.09	0.13 (I.10-4.42)	6.7	4.7
Neal & Latapie, 1973	е М	etiferus	80-83	0.1 hect. ponds	0.27	1.09 (0.83-16.2)	3.2	1.1
	P. 82	tecus	62-100	Ξ	0.08	0.47 (0.82-14.2)	12.6	1.6
Balazs <u>et al</u> ., 1973	P. 19	tecus	25	0.9 m <sup>2</sup> tanks	ł	4.03 (0.58-1.20)	NR <sup>f</sup> ,8	NRS
	년. 1	iponicus	25	-	<b>1.</b> 46	3.26 (1.50-6.44)	NRB	NRS
Shigeno <u>et al</u> ., 1972	빈	<u>iponicus</u>	25	1.7 m <sup>2</sup> tauks	5.30	8.68 (0.94-2.83)	$1.3^{\rm h}$	$1.0^{\rm h}$
		=	50	=	4.16	7.02 (1.73-8.11)	2.0 <sup>h</sup>	1.0 <sup>h</sup>
Deshimaru & Shigeno, 1972	빌	iponícus	60	1.7 m <sup>2</sup> tanks	12.71	15.13 (5.31-15.71)	1.6 <sup>h</sup>	1.3 <sup>h</sup>
Deshimaru & Kuroki, 1974a, b	년 1	<u>iponicus</u>	30	0.3 m <sup>2</sup> tanks	1.92	5.23 (0.87-2.49)	6.2	1.7
Andrews <u>et al</u> ., 1972	<u>Р</u> .	tiferus	56-70	0.8 m <sup>2</sup> tanks	0.07	0.28 (4-5.3)	NR	NR
Subrahmanyam & Oppenheimer, 1969	ਦ। ਚ	6 P S	42	l m <sup>2</sup> tanks	0.24	0.59 (0.61-1.28)	×100	5.5
Hysmith <u>et al</u> ., 1973	ы. В	, postlarval	30	0.2 m <sup>2</sup> tanks	0.14	3.15 (0.06-0.29)	6.9	1.68
	ଜା ଜା	, juvenile	32	0.2 m <sup>2</sup> tanks	4.41	16.16 (3.04-8.33)	21.78	1.48
Parker & Holcomb, 1974	ы Ч	, juv., Table 2	85-100	0.2 hect. ponds	0.27	1.08 (0.29-10.81)	3,8	J.f

	Table 17. (continu	led)		G	Dq	E) E4	1 10
<u>Reference</u>	Species <sup>a</sup>	LGP <sup>b</sup>	Inc		(M-FW) IH	IH	3
Caillouet <u>et al</u> ., 1973	P. duorarum	51-64	2 m <sup>2</sup> tanks	0.07	0.40 (1.3-6.6)	48.1 10 5	7.8
	-	98	.23 hect. po	nds V.Ið	( XN-TA ) 75 A	C'ot	T 0. T
Caillouet <u>et al</u> ., 1975b	<u>P. duorarum</u> Exp. I	63	2 m <sup>2</sup> tanks	0.05	0.49 (0.01-NR)	17.1	2.2
	" Exp. II	63	÷	0.10	0.31 (0.01-1.3)	17.9	5.6
F 74A, B, D, Purina "20"	P. duorarum	58-65	2 m <sup>2</sup> tanks	0.10	0.35 (0.01-1.35)	13	7
F 74A, B, D, wheat bran	=	=	Ŧ	0.08	0.27 (0.01-1.19)	16	6
F 74A, B, D, fungal feeds	=	2	=	0.04	0.20 (0.005-1.02)	50	13
F 74C, wheat bran	P. setiferus	60	=	0.15	0.15 (3.27-5.12)	8	œ
F 74C, fungal feeds	=	=	=	0.04	0.11 (3.27-4.53)	93	12
F 74C, fungal and bran	=	÷	F	0.11	0.15 (3.27-5.34)	12	æ

a Of the genus Penaeus.

b Length of growth period in days.

<sup>c</sup> Area of the bottom of containers in which growth took place.

calculated from tables of results given in the references cited. (IW-FW)= the average initial and final fresh weights Production of shrimp, in grams/m $^2$ /day. Range (LO-HI) is not necessarily the full range observed; rather, it is that observed for the high production value. σ

F E = feed efficiency=total dry weight of feed/final fresh weight of shrimp. Kange determined as for GMD. Θ

f NR= not reported or determinable from the reference cited.

<sup>E</sup> Excess food removed daily.

As footnote g, but only consumed food used in calculations, and weight of dead animals included. d

fact that our findings are the result of a very small-scale research effort.

14. Culture screening on delignified bagasse and straw

In our use of an agar plate method of screening, we attempted to circumvent the expense of instrument analysis and abbreviate the length of time necessary to determine ability of strains of fungi to convert test substrates into mycelial biomass. we diamor measure-me whom dottions obergonei abard who whom who have a flore whom in the strain of the str

fround to be the factor by windurspacific growth rate in redenerged culture can be related to colony radial growth rate on solid media. Had we done so, we would have arrived at a more accurate estimate of rate of increase in mass of tested fungi on each substrate than with only measurement of colony radial growth rate. Rather, we took into account hyphal diameter and hyphal density and used these to arrive at an indicator of fungal biomass produced. As Morrison and Righelato (1974) have found, hyphal density and hyphal diameter are important determinants of the width of the peripheral growth zone--it may be that refinement of methodology could yield an

estimate of specific growth rate using measures of these two characters rather than attempting the more exacting techniques of Trinci (1971).

There are objections which can be raised to the use of our methodology in this culture screen. Eslyn (1969), Nilsson (1974), Sharp and Eggins (1970) demonstrated that assay of fungal cellulolysis may yield different results when agar-plate methods and other more direct methods are used. These studies, however, did not involve attempts to measure biomass. Another important fault lies in the fact that differences among test fungi in depth of penetration into the agar were not measured (a few marked differences were noted), and surface density and hyphal diameter undoubtedly do not reflect submerged characters to the same extent among strains of fungi. Whether or not our method achieved approximation of conversion ability would be best determined by conducting correlation analysis of fermentation results (protein production, chitin production, available energy production) with our (or alternative)

computed indicators.

#### 15. Future directions

When we undertook this project, it was the feeling of some of those already involved in the artificial detrital feed research, and some of those who were to sponsor our research, that the problem to be treated was a rather simple one: just grow up some fungi on an inexpensive substrate, submit it to the young shrimp, and raise them to marketable size. After all, they were eating the same stuff in nature, weren't they, and growing just fine there? And fungi have been growing on lignocellulosic materials in nature for eons. All that would be necessary would be to bring the process into the laboratory and accelerate it.

Pernaissione consider most significant on furtiging in about the problem of developing a successful process for producing and utilizing natural food analogues is not as simple as it may seem on the surface. The series of steps in a functioning process is diagrammed in Fig. 12. As Calam (1969a) points out, research into particular fermentation processes has demonstrated "how stubborn and complex a fermentation projecticam be a fermentation projecticam be a fermentation studies of L. V. Sick and his colleagues (Sick and Baptist, 1973; Sick <u>et al.</u>, 1973) show that even inducing satisfactory ingestion rates can be a very complex problem. Regnault <u>et al.</u> (1975) should be consulted regarding problems of form of feed materials.

In order to successfully develop these steps in Fig. 12 so that they function as a working series, several factors or variables must be considered for each step. In many cases, the variables considered under a given step will have significant interacting effects on the functioning of that step, and to further complicate matters, variables which must be considered under one step may have distinct effects

ingus capable of converting

12. As a further example, suppose we were to identify a

a treated lignocellulosic substrate under a given set of fermentation conditions such that the fermented product contained a theoretically optimal amino acid-reservecarbohydrate-lipid-chitin balance, examining as many variables under each of the steps in Fig. 12 as possible. After having done all of this exhaustive work, our product might well turn out to be a dismal failure, returning us to our starting position, because of mycelial unpalatability or even toxicity to our target aquacultured organism; Nikolei (1961) found that unpalatability and/or toxicity appeared to be the causes of failures of his gall-fly larvae to grow well on some of his fungal feeds, and that this could be caused by strain of fungus and/or fungal culturing conditions!

With a little imagination and pessimism, one can conjure up several other discouraging scenarios. On the other hand, if the problem is dealt with intelligently, it should be amenable to solution. We would propose that success not be attempted in leaps and bounds (as it was in the present study--success under these conditions could only be the result of serendipity); rather, careful, thorough analysis of some of the most basic questions should be the primary effort. The three major goals of early work would be fungal strain and substrate selection, determination of optimal delignification method, and determination of optimal type of fermentation.

Fungal strain(s) which would serve well in the process must be capable of both efficient conversion of the substrate used and satisfactory support of shrimp growth. Therefore, we feel that the problem of strain selection should be dealt with by conducting a dual screen of as many strains as possible, examining concomitantly the degree of ability of the fungi for substrate conversion and acceptability and nutritive capacity of the mycelia for the shrimp. Those which ranked high in both categories would then be used in development of the fermentation and feed preparation processes. The thermophilic fungi make prime initial candidates for this survey (Barnes et al., 1972; Cooney and Emerson, 1964; Romanelli et al., 1975). Genetic engineering (Esser, 1974) and alteration of fermentation conditions could then be used

as tools in improvement of the fermentation product in terms of conversion efficiency and nutritive capacity. A listing of cultures of potentially biodeteriogenic fungi available at three major culture collections is given by Denizel <u>et al.</u> (1974).

Substrate selection would be based on the criteria of low acquisition cost and relative facility of conversion into fungal biomass. Substrates of low acquisition cost would be those agricultural by-products of high content of lignified cellulose and thereby little value as ruminant feed. They would have to be readily available in or near areas in which mariculture of shrimp has likelihood of commercial success in order that collection and transportation costs be minimal. They would have to be amenable to simple, low-cost delignification processes such as some of those which have been developed by the U. S. Environmental Protection Agency (Rogers and Spino, 1973) and other investigators (e.g. Bender <u>et al.</u>, 1970; Han <u>et al.</u>, 1975; Hartley <u>et al.</u>, 1974; Heany <u>et al.</u>, 1973; the work of Bavor (1974) may also be of interest here) in order to permit development of a fermentation or rumination process with high conversion rates.

Sugar cane bagasse is an example of an agricultural by-product which may meet the substrate-selection criteria. It is grown in large quantities in tropical climes where temperature conditions and land and labor costs are most favorable to penaeid shrimp mariculture operations. It is attractive as a substrate because it is available at centralized locations in the sugar processing plants (Callihan and Dunlap, 1969), and has limited value as feed, fuel, paper, or construction material (Srinivasan and Han, 1969). Simple delignification processes have been used on bagasse with success in increasing markedly its susceptibility to both bacterial and fungal degradation (from 15% to 70-80%) (Callihan and Dunlap, 1971; Dunlap, 1969; Garcia-Martinez <u>et al.</u>, 1974). (It must be noted here that Cruz <u>et al.</u> (1967) claimed to have produced fungalfermented bagasse of 28% crude protein <u>without</u> delignification and <u>without adding a</u> <u>nitrogen source</u> to the fermentation medium; their methods are cursorily reported,



The basic series of steps in a hypothetical functioning artificial detrital aquacultural process. Figure 12.
and their results are of questionable accuracy.) See Mandels <u>et al</u>. (1974) for a review of the susceptibility of cellulosic materials to fungal lysis.

The choice of type of fermentation process which is to be used should be dictated by potential costs. A comparison of two basic types of fermentation processes, submerged-culture and solid-state, indicate that the latter is best suited for artificial detrital feed production. By "solid-state" fermentations we simply mean those which require only moistening, rather than submerging of the substrates to be fermented. Hesseltine (1972) has recently discussed the use of solid-state fermentations, and laments the fact that in the West, deep-tank, or submerged-culture, fermentation methods have been developed and used to nearly the exclusion of solidstate methods (a strong indication of this is the fact that solid-state apparatus is not dealt with in the reviews of fermentation design given by Blakebrough (1969), Solomons (1969), or Steel and Miller (1970) ). Hesseltine lists advantages of solid-state fermentations; the most important and basic of these is the most obvious--solid-state fermentations require only small volumes of water. This removes the necessity for large vessels, and simplifies aeration methodology. It eliminates the need for supplies of large volumes of water and the problem of its disposal (if inorganic nutrients are supplied in the proper concentration, washing of the product of the fermentation should not be necessary). It provides conditions which in themselves inhibit bacterial contamination (the fungi are favored by low moisture levels relative to the bacteria (Hesseltine, 1972; Gray and Williams, 1971)). Harvesting of the product is clearly a very simple matter relative to the harvesting problems of submerged-culture fermentation; the finished product could conceivably be fed directly to the animals to be cultured.

Each of these advantages is a partial reason for the adaptability of solid-state fermentation to low-level technological operation (this aspect is discussed by N. J. Poole in a privately communicated manuscript which is now in preparation for publication--Poole and Smith, 1975; see also Imrie, 1975). As Brook <u>et al.</u> (1969) point out, solid-state fermentations are particularly well adapted to be carried out with simple apparatus at low cost, since this method of fermentation has been traditionally used in the production of fermented foods such as tempeh for over a thousand years (Gray, 1970; Hesseltine, 1965). What it amounts to is the production of a "vegetable cheese" (term from Brook <u>et al.</u>) with emphasis on production of mycelium. An example of a pilot-plant process similar to that which we envision for use in production of artificial detrital feeds is described for tempeh production by

Steinkraus et al. (1965) and for production of "mold bran" by Underkofler et al. (1947). licity of these processes permit them to be operated and managed by persons and the second statement with The sim al technical skill and training; this advantage has been recently recognized of mini anoglous.et al. (1976), who discuss the likelihood that successful mariculture . by Tch a will probably involve use of highly labor-intensive plans. That the of crusta sic substrates which we have recommended could be adapted to lignocellu ons of this kind is evident from the success achieved by Hesseltine et al. fermenta e production of secondary metabolites on forage materials (including oat (1968) in 1 g solid-state fungal fermentation, and from the preliminary results of straw) us il. (1972) with solid-state fungal fermentation of paper and of N. J. Poole Barnes et leagues (personal communication) with solid-state fungal fermentation of and his c 3ery, 1972). straw (see

uld seem that the potential for development of valuable information from ito artificial detrital feeds is strongly indicated. If enough support for nilar to ours is forthcoming, and if thorough analysis of the set of basic ivolved in developing a successful process of production and utilization of is is carried out (rather than hasty attempts at short-circuiting this al research), then we feel that it is likely that effective prototypal etrital feeds will be developed in the near future. straw) us Barnes <u>et</u> and his c straw (see It w research projects s problems these fee

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## 16. Range of applicability

As a final note, it should be pointed out that artificial detrital feeds of the kind discussed above are not limited in their potential applicability to the husbandry of shrimp. Several other detritivorous or omnivorous animals are aquacultured and commercially valuable (e.g. carp, mullet, tilapia--see Burdach <u>et al.</u>, 1972). These presumably (perhaps especially the mullets) have a digestive physiology geared to the types of feeds (microbial-detrital complexes) which would be produced by the methods which we propose. Feeds of this type might help change the fact that the <u>methods which we propose</u>. Feeds of this type might help change the fact that the <u>methods which we propose</u>. Feeds of this type might help change the fact that the <u>methods which we propose</u>. Feeds of this type might help change the fact that the <u>methods which we propose</u>. Feeds of this type might help change the fact that the <u>methods which we propose</u>. Feeds of this type might help change the fact that the <u>methods which we propose</u>. Feeds of this type might help change the fact that the <u>methods which we propose</u>. Feeds of this type might help change the fact that the <u>methods which we propose</u>. Feeds of this type might help change the fact that the <u>methods which we propose</u>. Feeds of this type might help change the fact that the <u>methods which we propose</u>. Feeds of this type might help change the fact that the <u>methods which we propose</u>. Feeds of this type might help change the fact that the <u>methods which we propose</u>. Feeds of the second second the second the

rearing of channel catfish. Sea lamprey larvae have been successfully raised using yeast as the only food source (Han son <u>et al.</u>, 1974). Even the apparently lucrative business of raising marine turtles (Anon., 1974) might benefit from application of this type of feed, since fungal feeds have been suggested for use with plant-eating livestock (Griffin <u>et al.</u>, 1974) and growth of pigs on fungal protein has been shown to be equal to that produced by soybean meal and fishmeal (Duthie, 1975).

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