

OPTIMIZATION OF ITACONIC ACID (IA) PRODUCTION USING ORGANIC ACID PRODUCING FUNGI ISOLATED FROM SOIL

¹Sadiq B.B, ³Aliyu A., ¹Amina A. G., ¹Tanko N.D., ²Nasiru S., ¹Zulaihat L., ²Bala M.

Department of Biochemistry, Bayero University, Kano
 Department of Biochemistry, Federal University, Dutse
 Department of Biochemistry, Aliko Dangote University of Science and Technology Wudil, Kano

*Corresponding Author e-mail: sadiqbabuga63@gmail.com

Abstract: Microbial production of organic acids has been incessantly escalating over conventional chemical methods due to several advantages including enantio-selectivity, high purity, less environmental pollution and cost effectiveness. Itaconic acid is used as a platform chemical for the production of various value-added chemicals such as poly-itaconic acid, resins biofuel components, ionomer cements. This research work aimed to isolate Aspergillus terreus producing itaconic acid from soil and optimize the maximum yield conditions. Aspergillus terreus was isolated, screened and identified morphologically and molecularly and for itaconic acid production. The itaconic acid produced was confirmed by bromocresol purple and measured by UV spectrophotometer in which the optical density was taken at 385nm. Itaconic acid was optimized initially by one factor at a time (OFAT) and then response surface methodology (RSM) using Central Composite Design (CCD). Incubation time, substrate concentration, pH and temperature were optimized and obtained the maximum yield of 22.601mg/ml itaconic acid at incubation time of 96hrs, substrate concentration 10% of molasses, pH of 4.0 and temperature of 34.5°C with a significant quadratic model of P- value less than 0.05. The response surface plots (3D and contour) presented that the interactions between the parameters were significant to itaconic acid production. Based on this research Aspergillus terreus has a good potential influence for microbial production of itaconic acid using molasses as substrate.

Key words: Fermentation, itaconic acid, Aspergillus terreus, optimization, molasses

INTRODUCTION

of limited resources have led to the development producers (Becker et al. 2015). commodity chemicals for construction industries as well as for the various applications (El-Imam et al. 2014). commercially (Rajendra et al. 2017).

chemicals. and coatings, Ustilago zeae, U. maydis, Candida sp., and biodegradable plastic.

about 80,000 tons is currently met by fungal Increasing environmental concern and depletion fermentation relying on natural Aspergillus spp.

of microbial approaches for the production of Itaconic acid is used as a platform chemical for the sustainable production of various value-added chemicals such as development (Aliyu et al. 2022). The advances poly-itaconic acid, resins biofuel components, in microbiology and fermentation technology ionomer cements etc. Itaconic acid and its derivatives have led to the development of eco-friendly have wide applications in the textile, chemical and processes replacing some of the conventional pharmaceutical industries. The depletion of fossil chemical methods. Microbial production of fuels and the need for sustainable development organic acids has been incessantly escalating require that fermentative itaconic acid production over conventional chemical methods due to replace petroleum-based methods of itaconic acid several advantages including enantioselectivity, production. Various microorganisms have been high purity, less environmental pollution and employed in itaconic acid fermentations, with the cost effectiveness (Aliyu et al. 2022; Mari et al. most prolific producer being Aspergillus terreus. 2022). The increasing knowledge on metabolism Over 80 g/L itaconic acid has been produced in and pathway regulation of industrially relevant fermentations using glucose. However, there is an organisms has already proven to be invaluable increasing interest in the utilization of lignocellulosic for generating rational strain development with materials for itaconic acid production due to the primarily industrial performances. In recent concern of food security. The current industrial times, microbial production of building block applications of itaconic acid and its potential use as a chemicals is progressively expanding its market drop-in or novel substitute monomer to replace for the production of succinic, lactic, citric, petroleum-based chemicals were also extensively itaconic, gluconic, lacto bionic acids, etc. explored. Recent trends in itaconic acid research Organic acids are utilized directly or indirectly summarized that itaconic acid can be produced cost in a wide range of applications including food, effectively from sustainable raw materials and have healthcare, cosmetics, textile, solvents, and the potential to replace petro-based chemicals in

manufacturing of biodegradable packaging In contrast with citric, gluconic, and lactic acids, materials. Organic acids with wide applications itaconic acid is used exclusively in non-food in various fields are made from living cells applications. Its primary application is in the polymer industry where it is employed as a co-monomer at a Itaconic acid is an important component in the level of 1-5% for certain products. Itaconic acid is chemical industry. Itaconic acid (IA) and its also important as an ingredient for the manufacture of derivatives have broad application spectrum in synthetic fibers, coatings, adhesives, thickeners, and pharmaceutical binders. The market volume has been estimated to be industries. It is used as building block for acrylic about 15,000 metric tons per year and is expected to plastics, acrylic latexes, anti-scaling agents and grow if the selling price (estimated to be about US\$4 super absorbents (Steiger et al. 2013). It also has per kg) can be reduced (Willke and Vorlop 2001). To other applications in food packaging, detergents, date very little research has been directed at the pharmaceuticals, improvement of itaconic acid production. In contrast, agriculture, emulsifiers, herbicides and printing there has been a larger research effort directed at chemicals. Many microorganisms, such as lactic acid production to feed the market for

Rhodotorula sp. have been reported to produce Itaconic acid was historically produced by various itaconic acid (Kawamura et al. 1981). However, chemical methods which includes Destructive Aspergillus terreus is a preferred source in distillation of citric acid, and this was the main commercial production of itaconic acid up to 80 method of producing it prior to the 1960s; Oxidation g/L (Steiger et al. 2013). The annual market of of isoprene etc. None of these (or other) processes

and itaconic acid (IA) is now almost entirely the attendant deleterious environmental effects. produced by fermentation of sugars by MATERIALS AND METHOD Aspergillus terreus (Tsao 1990). The Northern Sampling of the Soil Regional Research Laboratory (NRRL) of the The soil sample was collected at Bayero University went on to become the most published strain.

The initial industrial production of itaconic acid Medium Preparation market volume for itaconic acid (IA) is about of fungi. 80,000 tons per year in 2005 and the selling Isolation of Aspergillus terreus from the Soil price is \$2.00 per kg (Okabe et al., 2009), and Sample processing and production methods.

production. Bio-based itaconic successfully produced from renewable biomass. medium. The high request for bio-based materials Morphological Identification of Aspergillus terreus anticipates a further growth of 60% of the Staining Reaction itaconic acid (IA) world market, which is The pure fungus of Aspergillus terreus grown as chemical which has a wide range of actual and (Shalini et al. 2014).

compete favourably with the fermentative wide range of petroleum based chemicals, e.g. acrylic production process (Willke and Vorlop 2001), acid, which will reduce dependence on petroleum and

United States Department of Agriculture Kano, old campus (Latitude of 11.9742°N and (USDA) screened several wild type strains and Longitude of 8.4684°E) Gwale LGA, Kano Nigeria identified Aspergillus terreus NRRL 1960, as and collected from a dark loamy soil and transferred the most prolific itaconic acid (IA) producing safely to the laboratory. Exactly 1 g of the soil strain (Lockwood and Reeves 1945) which then sample was weighted and diluted with 9ml distilled water using test tubes.

used a chemical approach that is the pyrolysis of The primary medium used for isolation of fungi was citric acid to itaconic anhydride, followed by the potato dextrose agar (3.9 g of agar was dissolved in hydrolysis of the anhydride which leads to the 100 ml of distilled water). The medium was production of nonrenewable chemicals. The autoclaved at 121°C and 15 lb pressure for 15 min. chemical synthesis of itaconic acid (IA) is not The medium was poured on Petri plates in sterilized very effective. Thus, it is estimated that the laminar flow to avoid contamination for the growth

there is an expectation for a market demand From the collected soil sample (Soil dilution method: increase if the selling price can be reduced. Hawaz et al., 2023) diluted with 1 g soil is in 10ml of Global carbon emissions as a result of sterile distilled water. Exactly 9 ml of distilled water petroleum-based processing and products are was taken to 5 different test tubes and 1 ml of the soil forcing industries to look for alternative liquid suspension was serially transferred to each 9 ml distilled water containing test tubes. 1 ml of Production of itaconic acid by fermentation suspension from each 5 different test tubes after needs significant improvements which will dilution was added to sterile Petri plates in duplicates increase the productivity and reduction costs of containing sterile Potato Dextrose Agar (PDA) acid medium. The medium was autoclaved at 121°C and production promises to be an attractive 15 lb pressure for 15 min. The plates were incubated alternative for the chemicals industry to replace at 28°C for 5- 7 days. The growth of separate petroleum-based chemicals synthesis. Many colonies was observed. A greater number of species chemicals such as succinic acid, 1, 3- were isolated most of the fungus speculates heavily. propanediol and ethanol which were hitherto Pure culture was done using Petri plates in duplicates made from petroleum refining are now being for each colony containing fresh agar of PDA

predicted to surpass 216 million USD in the year metallic brown colonies or different colours. From 2020. According to the annual forecast, market the bottom, the colonies confer on the medium a dark is predicted to exceed 410,000 tons of itaconic brown color. The isolated fungi were identified to the acid by the year 2020 (Choi et al. 2015). genus level and to the species when possible on the Itaconic Acid (IA) is an important platform basis of macro morphology and micro morphology

potential applications. It can be used to replace a The macro morphological is by observing the colony

growth, topography (flat, heaped, regularly or (Aliyu et al. 2022). irregularly folded), texture (yeast like, powdery, One Factor at a Time (OFAT) for Itaconic Acid velvety cottony), or (pigmentation and reverse pigmentation).

identifications. The identified /spores/conidia and cytoplasm were stained by taken at 385nm using a UV Spectrophotometer. The stained specimen (Aspergillus duplicate, agent. observed under the terreus) microscope (Magnus MLXi plus) under $10X \times 40X$ magnification.

Pretreatment of Molasses

cane molasses was diluted upto 100ml with taken at 385 nm using a UV Spectrophotometer. distilled water. Then 5ml of the acid was added For optimum pH determination, six Erlenmeyer (Shazia *et al.*, 2015).

Determination of Optimum Growth Conditions for Itaconic Acid (IA) Production

concentration, pH and Temperature) were using a UV Spectrophotometer. considered for optimum growth of the fungal For optimum temperature

features (color, shape, size and hyphae). The response surface methodology (RSM) by using colonies were examined for slow or for rapid Design Expert version 6.6.0 optimization software

surface (IA) Yield

For optimum incubation time determination, six The micro morphological is by a compound Erlenmeyer flasks having 100 ml of Czapec Dox microscope with a digital camera using a lacto broth were prepared in duplicate, and their incubation phenol cotton blue-stained slide mounted with a time were adjusted at 24 hrs, 48 hrs, 72 hrs, 96 hrs, small portion of the mycelium (Gaddeyya et al. 120 hrs and 144 hrs which were then autoclaved. The 2012). The Hyphae, macro conidia, micro Erlenmeyer flasks were properly inoculated with conidia, chlamydospores and other special freshly prepared spores suspension (inoculum size) of fungal structure, characteristics using suitable fungal isolates and incubated under proper conditions media, slide cultures and the most updated keys at 35°C, pH of 5.0 and substrate concentration fungi (molasses) of 3% in an incubator shaker at constant confirmed with microbial expert. The fungal rpm (200) were used to investigate the influence of propagules were coloured. The coloured mycelia itaconic acid (IA) production, their absorbance were

using Lacto phenol and cotton blue. Cotton blue For determining the optimum percentage of substrate were stained cytoplasm and results in light blue concentration (molasses), six Erlenmeyer flasks background. Lacto phenol acts as a cleaning having 100 ml of Czapec Dox broth were prepared in their and percentage substrate light concentration (molasses) were adjusted to 2%, 4%, for 6%, 8%, 10% and 12% which were then autoclaved. identification and microphotograph was taken The Erlenmeyer flasks were properly inoculated with freshly prepared spores suspension (inoculum size) of fungal isolates and incubated under proper conditions Molasses are thick brown sweet liquid that is at temperature of 35°C, pH of 5.0 and incubation time made from raw sugar. Cane molasses was of 120 hrs in an incubator shaker at constant rpm pretreated with acid (H₂SO₄). 2.0 N of the acid (200) were used to investigate the influence of was used for molasses pretreatment. 15ml of the itaconic acid (IA) production, their absorbance were

and placed in a water bath at 90±2°C for 1 hour. flasks having 100 ml of Czapec Dox broth were After cooling at room temperature, the medium prepared in duplicate, and their pH were adjusted to was neutralized with lime (CaO) and left to 3.0, 3.5, 4.0, 5.0, 6.0 and 7.0 which were then stand overnight. Two layers were formed, the autoclaved. The Erlenmeyer flasks were inoculated upper shiny black and lower yellowish brown with freshly prepared spores suspension (inoculum due to the presence of trace metals. The clear size) of fungal isolates and incubated under proper supernatants were diluted to desired sugar level conditions at temperature of 35°C, incubation time of 120hrs and substrate concentration (molasses) of 3% in an incubator shaker at constant rpm (200) were used to investigate the influence of itaconic acid (IA) Four parameters (Incubation time, Substrate production, their absorbance were taken at 385nm

determination, isolates for the production of itaconic acid (IA). Erlenmeyer flasks having 100ml of Czapec Dox One factor at a time (OFAT) method of broth were prepared in duplicate, and their optimization was performed first before using temperature were adjusted to 30°C, 32°C, 35°C, 37°C,

proper conditions at incubation time of 120hrs, the model. substrate concentration (molasses) of 3% and Data Analysis pH of 5.0 in an incubator shaker at constant rpm The average data and standard deviations were itaconic acid (IA) production, their absorbance run using Microsoft Excel (Office, 2019). The taken at 385nm using Spectrophotometer (Meena et al., 2010).

Itaconic Acid (IA) Yield

In general, the change in culture conditions considered significant and vice versa. greatly influenced the production ability of RESULTS AND DISCUSSION itaconic acid (IA) synthesis. The statistical base Morphological Identification of Aspergillus terreus effect incubation time, (Table 1), this resulted to thirty experimental Aspergillus terreus respectively. The macroscopic runs from the software Design expert, version and microscopic characteristics of AT8 (Aspergillus 6.0.6. The modeling and data analysis were terreus) is presented in Table 2 and Table 3 6.0.6 (Aliyu et al. 2022).

Table 1: Factors for RSM Experimental Design.

Indicator	Factor	Low level	High level
A	Incubation time	72	120
В	Substrate conc.	8	12
C	pН	3	5
D	Temperature	32	37

Validation of the Second Order Polynomial Model

The second order polynomial model obtained from RSM was validated by conducting a series of experiments selected at random from the design in Table 5. The experiments were done by choosing random values of parameters within

40°C and 42°C which were then autoclaved. The the optimized levels as presented in Table 5 below. Erlenmeyer flasks were properly inoculated with The experimental output was then compared to the freshly prepared spores suspension (inoculum values predicted by the second order model obtained size) of fungal isolates and incubated under from CCD, to estimate the fitness and goodness of

(200) were used to investigate the influence of obtained from the duplicate of experiments for each a UV standard deviation for each value was 5% analysis of variance (ANOVA) was done sing Design-Expert Response Surface Methodology (RSM) for software 6.0.6. A confidence level of 95% was used in this study. Any p-value less than 0.05 was

optimization was used to study the influential The results obtained from the morphological substrate (macroscopic and microscopic) analysis of isolated concentration (molasses), pH and temperature organisms (AT2, AT4, AT6 and AT8) have shown on itaconic acid (IA) yield using Central variation in colony colours, reverse colours, margin Composite Design (CCD). The optimized covering and growth level. The macroscopic and conditions of parameters were taken as microscopic characteristics of isolates show that independent variables and the itaconic acid (IA) AT2, AT4, AT6 and AT8 are Aspergillus niger, yield was chosen as the dependent variables Aspergillus fumigatus, Penicillium crysogenum and

performed sing Design expert software, version Table 2: Screening of isolates for the production of itaconic acid (IA).

Isolates Code	AT2	AT4	AT6	AT8
Indicator (BCP)	+	_	_	+

KEY: AT2 = Aspergillus niger, <math>AT4 = Aspergillusfumigatus, AT6 = Penicillium crysogenum, AT8 = Aspergillus terreus, BCP = Bromocresol purple.

Table 3: Morphological Characteristics of Aspergillus terreus

Characteristics	Aspergillus terreus		
Hyphae	Septate and hyaline		
Conidiospore	Smooth and hyaline		
Conidia heads	Compact		
Vesicle	Biseriate spherical		
Shape	Glubose		

Table 4: Microscopic Characteristics of Aspergillus terreus

Characteristics	Aspergillus terreus		
Surface colour	Light yellow to metallic dark brown colour.		
Growth	Slow.		
Margin covering	Half to one-third		
Reverse of the colony	Appears metallic brown		

Molecular Identification of Isolated Fungi

The results for molecular identification of the fungal isolate are presented in Figure 4.1 and 4.2. Figure 4.1 presented gel electrophoresis of the 18S ribosomal RNA (18SrRNA) gene result for AT8 showing 600 base pairs on the DNA molecular weight ladder, the sequence for the 18S rRNA of the isolate confirmed the identity of the isolate (AT8) as Aspergillus terreus with 99% identity with Aspergillus terreus MW881456 with accession number of OP8866158 after blasting in NCBI.

Production of Itaconic Acid (IA) using One Factor at a Time (OFAT)

Figure 1, 2, 3, and 4 presented the optimum production of itaconic acid results at different parameters (incubation time, substrate concentration, pH, and temperature) using one factor at a time (OFAT) technique.

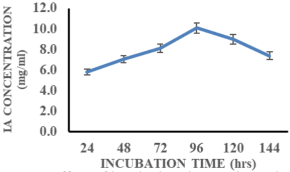


Figure 1: Effect of incubation time on itaconic acid production at constant substrate concentration (3%), pH (5) and temperature (35°C) using one factor at a time (OFAT).

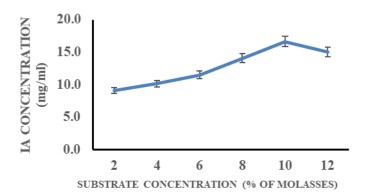


Figure 2: Effect of substrate concentration on itaconic acid production at constant incubation time (96hrs), pH (5) and temperature (35°C) using one factor at a time (OFAT).

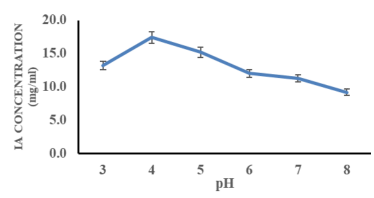


Figure 3: Effect of pH on itaconic acid production at constant incubation time (96hrs), substrate concentration (10%) and temperature (35°C) using one factor at a time (OFAT).

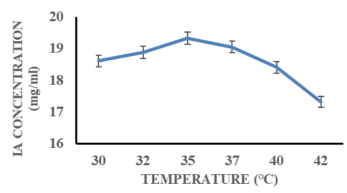


Figure 4: Effect of temperature on itaconic acid (IA) production at constant incubation time (96hrs), substrate concentration (10%) and pH (4.0) using one factor at a time (OFAT).

Incubation Time Optimization

Figure 1 presented the result of gradual increase in itaconic acid (IA) yield with increasing incubation time from 24hrs to 96hrs and later declined at 120hrs and 144hrs. The optimum itaconic acid (IA) yield (10.09mg/ml) was obtained at 96hrs

Substrate Concentration Optimization Substrate concentration percentage of molasses has effect in itaconic acid (IA) yield in which it was increased with increase in substrate concentration from 2% to 10%. The optimum itaconic acid (IA) yield (16.65mg/ml) was obtained at 10%. In effect, it later declined at 12% of the substrate concentration. Figure 2 presented the result of gradual increase of itaconic acid (IA) yield by effect of substrate concentration percentage.

pH Optimization

Increase in itaconic acid (IA) yield was observed when pH increased from 3.0 to 4.0 and AC, AD, BC, BD, CD are the intermassively declined from pH 5.0 to pH 8.0. The ptimum yield of itaconic acid (IA) was achieved at pH of 4.0 (17.415mg/ml). The result of itaconic acid (IA) yield with effect in pH is presented in Figure 3.

Temperature Optimization

Increase in itaconic acid (IA) yield was observed with increase in temperature from 30°C to 35°C, the optimum itaconic acid (IA) yield (19.05mg/ml) was obtained at the temperature of 35°C and then later declined from 37°C. Figure 4 presented the effect of temperature on itaconic acid (IA) yield.

Optimization using Response Surface Methodology (RSM)

Results obtained for actual itaconic acid (IA) concentration at different conditions of incubation time, substrate concentration, pH and temperature from response surface methodology (RSM) was presented in Table 5, the results of the responses obtained for each experimental run and the predicted responses were closer to each other.

Model F-value of 19.96 implies the model is significant. There is only a 0.01% chance that a "Model F-Value" this large could occur due to

noise. Values of "Prob > F" less than 0.0500 indicate model terms are significant. In this case A, A^2 , B^2 , C^2 , D^2 , AC, BC, BD are significant model terms. Values greater than 0.1000 indicate the model terms are not significant. If there are many insignificant model terms (not counting those required to support hierarchy), model reduction may improve your model. The "Lack of Fit F-value" of 0.90 implies the Lack of Fit is not significant relative to the pure error. There is a 58.92% chance that a "Lack of Fit F-value" this large could occur due to noise. Regression analysis produced the following second order polynomial fit with a satisfactory coefficient of determination ($R^2 = 0.9490$).

Itaconic acid concentratin= +21.91+1.53A+0.36B +0.37C +0.21D -2.30A² -1.90B² -3.08C² -0.65D² -0.56AB+0.74AC-0.57AD+0.82BC-1.22BD+0.46CDEquation ii

Where A, B, C and D are incubation time, substrate concentration, pH and temperature respectively. AB, AC, AD, BC, BD, CD are the interactions, and A², B², C², D² are the quadratic terms.

Table 5: Actual and predicted itaconic acid yield at different condition of incubation time, substrate concentration, pH and temperature.

Run	Incubation time (hrs)	Substrate concentration (ml)	рН	Temperature (degree)	Experimental value (mg/ml)	Predicted valu (mg/ml)
1	96.00	6.00	4.00	34.50	14.25	11.18
2	96.00	14.00	4.00	34.50	14.54	15.03
3	120.00	8.00	5.00	37.00	16.25	13.81
4	72.00	12.00	3.00	32.00	13.02	15.41
5	96.00	10.00	6.00	34.50	11.12	7.88
6	96.00	10.00	4.00	34.50	21.35	21.91
7	72.00	12.00	5.00	32.00	14.32	13.80
8	72.00	12.00	3.00	37.00	13.55	18.36
9	120.00	12.00	3.00	32.00	16.75	14.25
10	72.00	8.00	5.00	37.00	13.15	15.81
11	120.00	12.00	5.00	32.00	17.00	12.01
12	120.00	8.00	3.00	37.00	16.99	11.32
13	96.00	10.00	4.00	29.50	19.67	12.79
14	96.00	10.00	4.00	34.50	19.85	17.32
15	72.00	8.00	3.00	32.00	11.75	13.84
16	120.00	12.00	5.00	37.00	17.25	16.12
17	96.00	10.00	4.00	34.50	21.85	19.63
18	96.00	10.00	4.00	34.50	22.55	21.76
19	72.00	12.00	5.00	37.00	13.25	13.60
20	72.00	8.00	3.00	37.00	13.75	15.02
21	96.00	10.00	2.00	34.50	8.23	8.84
22	96.00	10.00	4.00	39.50	19.12	16.34
23	120.00	8.00	5.00	32.00	14.85	18.90
24	96.00	10.00	4.00	34.50	22.45	19.73
25	120.00	8.00	3.00	32.00	13.75	16.91
26	144.00	10.00	4.00	34.50	16.12	19.91
27	96.00	10.00	4.00	34.50	21.27	21.91
28	120.00	12.00	3.00	37.00	10.33	13.91
29	48.00	10.00	4.00	34.50	9.44	12.91
30	72.00	8.00	5.00	32.00	7.00	12.91

Table 6: Analysis of Itaconic Acid Yield from Quadratic Model Analysis of Variance.

Source	Sum of Square	Mean Square	F- Value	P > F	
Model	521.85	37.28	19.96	< 0.0001	Significant
A	56.24	56.24	30.11	< 0.0001	_
В	3.05	3.05	1.63	0.2205	
C	3.35	3.35	1.79	0.2008	
D	1.03	1.03	0.55	0.4685	
A2	145.60	145.60	77.95	< 0.0001	
B2	99.04	99.04	53.02	< 0.0001	
C2	260.23	260.23	139.32	< 0.0001	
D2	11.60	11.60	6.21	0.0249	
AB	5.06	5.06	2.71	0.1205	
AC	8.82	8.82	4.72	0.0462	
AD	5.22	5.22	2.80	0.1153	
BC	10.82	10.82	5.79	0.0294	
BD	23.77	23.77	12.72	0.0028	
CD	3.40	3.40	1.82	0.1970	
Lack of Fit	17.98	1.80	0.90	0.5892	Not
					Significant
R-Squared	0.9490	Adj R-	0.9015	Pred R-	0.7854
		Squared		Squared	

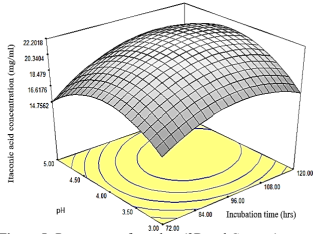


Figure 5: Response surface plots (3D and Contour) presenting the interaction between Incubation time and pH affecting itaconic acid (IA) production.

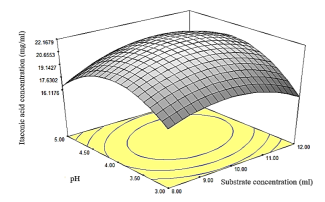


Figure 6: Response surface plots (3D and Contour) presenting the interaction between Substrate concentration and pH affecting itaconic acid (IA) production.

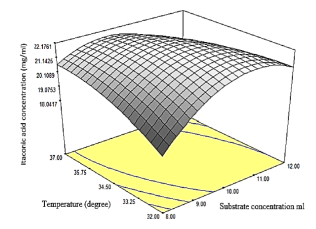


Figure 7: Response surface plots (3D and Contour) presenting the interaction between Substrate concentration and Temperature affecting itaconic acid (IA) production.

3D-Response Surface Plots Representing the Interaction between the Variables

Interaction among the various factors and the determination of optimum condition for maximum itaconic acid (IA) production were studied by plotting three-dimensional (3D) — and contour response surface plot, as presented in Figure 5, 6 and 7.

The results obtained shown the parabola shape of the 3D plot and the circular shape of contour plots indicated the interaction between incubation time and pH was significant, keeping substrate concentration and temperature constant. Figure 5 presented the results of response surface (3D and contour) obtained for the interaction between the incubation time and pH.

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The results obtained shown the parabola shape of the 3D plot and the circular shape of contour plots indicated the interaction between substrate concentration and temperature was significant, keeping incubation time and pH constant. Figure 7 presented the results of response surface (3D and contour) obtained for the interaction between the substrate concentration and temperature.

Validation of the Second Order Polynomial Model between the Experimental and Predicted Value of Itaconic Acid (IA)

The results obtained indicated that there was very good correlation between experimental and predicted values and in turn proved the validity of the models. The observed values of itaconic acid (IA) yield were compared with the values of predicted by the second order model. Table 7 presented the result of validation runs with observed and predicted values.

Table 7: Validation of the Second Order Polynomial Model between Experimental and Predicted Value of Itaconic Acid (IA) yield.

racome racia (11) freia.						
Ru n	Incubation time (hrs)	Substrate conc. (%)	pН	Temp (°C)	Experimenta l value (mg/ml)	Predicted value (mg/ml)
1	103.78	10.11	4.1 1	34.36	22.481	22.204
2	96.00	10.00	4.0 0	34.50	22.531	24.000
3	96.00	10.00	4.0	34.50	22.601	22.550

DISCUSSION

In this research study, four different fungal isolates were isolated from soil and identified morphologically and molecularly. AT8 (Aspergillus terreus) show the highest itaconic acid production capability among the isolates (Helia and Wan 2015) and (Meena et al. 2010). Molasses is used as source of glucose which is a very convenient raw material for itaconic acid production (Lockwood and Reeves 1945). DNA sequencing and molecular identification from Genbank identified isolate AT8 as new strain of Aspergillus terreus with closet similarity of 98% identity with Aspergillus terreus MW881456. Aspergillus terreus assigned the accession number of OP866152. Sequence alignment and evolutionary history was gathered for plotting phylogenetic tree.

Itaconic acid was produced under the influence of physicochemical parameters which were determined to have high yield by optimizing them with conventional technique, one factor at a time (OFAT) and response surface methodology (RSM) (Sadiq *et al.* 2017). The parameters were maintained significantly for better fermentation yield.

Effect of incubation time on itaconic acid (IA) was observed by varying time from 24hrs to 144hrs and keeping all other variables constant. The optimum yield of itaconic acid (IA) was determined at 96hrs of incubation and declined with increase incubation time as presented in Figure 4.3, this was in line with the study done by Linda (2021) by determining the optimum incubation time of the fungal isolates. The itaconic acid yield declined due to the relationship between the *Aspergillus terreus* and sugar contents in the medium and the fungal growth curve. Microbial production of metabolites usually starts after a lag

onset of stationary phase or late.

Effect of substrate concentration on itaconic acid (IA) was observed and the optimum itaconic acid produced (16.65mg/ml) was The obtained at 10% concentration of molasses. The Aspergillus terreus cell number increased exponentially which could provide the maximal conversion of substrate to itaconic acid was found at 10% this was related with the work of produced at molasses concentration above 10% secondary byproducts that limit itaconic acid (IA) productivity in accordance to the research done by Hawaz et al. (2023).

Effect of medium pH was revealed to have the itaconic acid (IA) yield 17.415mg/ml at pH of 4.0, which was found to be similar with the work of EL Imam et al. (2013) and Sudarkodi et al. (2012) who reported a maximum yield of itaconic acid (IA) found to be at pH 4. The enzymatic reactions in the utilization of energy are regulated by pH. Basically, the impact of low pH is associated with the activity of enzymes taking part in the biosynthesis of itaconic acid (IA) and subsequent transfer mechanism to the extracellular space/out of the cell. This is related to the finding of Peter et al. (2019) reported that itaconic acid (IA) generation by filamentous fungi such as Aspergillus terreus favors lower pH conditions and it has been argued that besides enabling the appropriate growth of Aspergillus terreus, such a fermentation environment can be useful for suppressing the formation of by-products that would lower the final itaconic acid (IA) yield and productivity.

Effect of temperature on itaconic acid (IA) was also observed and the maximum itaconic acid 35°C. After the optimum temperature the overall al. (2010). This high value of cell death error sum of squares (Mohamed et al. 2013). increases with increase in temperature, than the

phase of one day and reaches maximum at the growth rate. Hence the overall growth rates rapidly declined above the optimal temperature. Apart from this, the product inhibition effect is also more at higher temperatures than at lower temperatures.

results obtained from the preliminary optimization using one factor at a time (OFAT) experiment were then applied to response surface methodology (RSM) modelling. 30 experimental runs using central composite design (CCD) were design as presented in Table 5, the experimental yield was Meena et al. (2010). Low itaconic acid (IA) recorded and relatively closed to the predicted value presented in Table 5. The statistical analysis for and this may occur due to the formation of significances of all factors was described by analysis of variance (ANOVA) in Table 6. Based on the result obtained, the model of analysis was confirmed significant and highly reliable at P-value (< 0.0001) less than 0.05, and the no significant of lack of fitness indicated that the model was excellent fitted with no significant noise, the R² and adjusted R² value were all closed to 1 (0.9490 and 9015 respectively) showing the goodness of the model (Table 6). Additionally, the significant of the interaction between incubation time and pH (AC), substrate concentration and pH (BC) and that of substrate concentration and temperature were presented in ANOVA result (Table 6) with P-value 0.0462, 0.0294 and 0.0028 respectively. Furthermore, the response surface plots (3D and contour) reveal that the interaction between the factors (incubation time, substrate concentration, pH and temperature) were all significant to itaconic acid (IA) production as presented in Figure 5, 6, and 7. This shown that the itaconic acid (IA) production was dependent on the four parameters optimized.

> Validation of the second order polynomial model confirmed the optimum itaconic acid (IA) yield of (22.601mg/ml) and indicated linear interaction and quadratic effect of variables on itaconic acid production. Therefore, the developed model is considered reliable.

(IA) production of 19.05mg/ml was obtained at The experimental and the predicted value were relatively closed indicating the relative fitness of growth rate began to fall due to increase in rate experimental model. Moreover, this shows that of microbial death, as the death rate is also a extraneous factor terms in a derived model equation function of temperature as reported by Meena et will affect in some reduction in the calculation of the

CONCLUSION

According to the results obtained from this study, the isolated fungal species (Aspergillus terreus) has the potential ability of utilizing molasses as substrate for itaconic acid (IA) production. The study revealed that itaconic acid (IA) could be effectively yielded by adjusting incubation duration, and substrate concentration.

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