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# Reproductive characteristics of the sago strain gourami (*Osphronemus goramy* Lacepède, 1801): basic knowledge for a future hatchery strategy development

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# Abstract

**Background:** The sago strain gourami (*Osphronemus goramy*) has been approved in 2018 as a candidate for freshwater aquaculture in Indonesia. However, information on the species' reproduction is minimal. This study analyzed the reproduction characteristics of the sago strain gourami broodfish to provide basic knowledge for a hatchery development strategy in the future.

# Methods:

A total of 20 female and male broodfish sago strain gourami had matured oocytes were measured for body weight and length, and were evaluated for their reproductive characteristics. Breeding fish were spawned naturally in a 2×1×0.6 m concrete pond with a male-female sex ratio of 1:1. Egg weight and diameter were measured in 25 eggs per female using ACIS AD- 600i scales with 0.01 g accuracy and a microscope (Labo model L-711) using software camera 3. Semen was collected using plastic syringes in 3 mL aliquots, then placed in an insulated ice-cooled container, and analyzed within 2 hours of collection.

# **Results:**

\_Average weights of female and male broodfish before spawning were 2180±159.78 g and 3060±134.99 g. The relative fecundity and egg diameter were 1029±36 eggs kg<sup>-1</sup> and 2.42±0.05 mm. The hatching rate and embryo survival to eyed-egg stage were 76.40±2.27% and 94.76±0.42% Sperm characteristics showed that volume was 0.60±0.12 ml kg<sup>-1</sup> and percentage motile was 70.04±2.27%. Female broodfish weight after spawning had strongly corelated with the female broodfish weight after spawned ( $r^2 = 0.999$ ) and absolute fecundity had also strongly corelated with

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female broodfish before spawning ( $r^2$  = 0.921). While, the parameter of sperm concentration has a moderately corelated with the sperm motility ( $r^2$  = 0.556) and duration of sperm motility ( $r^2$  = 0.502).

# Conclusions:

The sage strain gourami broodfish has suitable reproductive characteristics for the development of future hatcheries. Successful natural spawning should be followed by larval weaning and feeding technology to increase growth and survival.

Keywords: Aquaculture, giant gourami, broodfish, egg, sperm, hatchery performance.

# Introduction

Freshwater aquaculture practiced in inland waters such as lakes, rivers, reservoir, floodplains and oxbow lakes, and freshwater ponds, has expanded during the last decades in Indonesia<sup>1,2,3,4,5</sup>. Approximately 77.57% of fish produced in freshwater aquaculture in\_Indonesia are sourced from freshwater ponds and inland waters<sup>6</sup>. However, its development depends upon many factors, such as fish species, aquaculture systems, water depletion, fish diseases, farmers'\_knowledge and aquaculture practices<sup>7,4,8,9,10</sup>.

Freshwater aquaculture is one of the fastest growing aquacultures in Indonesia, with more than 3,378,298.92 metric tons produced in 2018<sup>6,11,12</sup>. Nile tilapia (*Oreochromis niloticus*) contributed 37.93% of the total aquaculture production, African catfish (*Clarias gariepinus*\_33.35%, *Pangasius* catfish (*Pangasius hypophthalmus* 12.38%, common carp (*Cyprinus carpio*) 9.28% and giant gourami (*Osphronemus goramy*) 6.96% <sup>13,14,15</sup>.

Indonesian giant gourami strains\_include the local "tambago", "palapa", "soang", "galunggung" and "blusafir" strains, which have been grown semi-intensively in small-scale farms for decades<sup>8,14,13,16</sup>. However, they have not been able to contribute majorly to- freshwater aquaculture production in Indonesia.Therefore, worthy this a concern to be development of sago strain gourami was living a limited in West Sumatera Province of Indonesia<sup>17,18</sup>. The sago strain gourami is considered a source for nutritional and food security among many freshwater fishes' communities in Indonesia.

The sago strain gourami was approved as a candidate for freshwater aquaculture in 2018 (Decree of the Ministry of Marine and Fisheries, Republic of Indonesia No.56/KEPMEN-KP/2018)<sup>19</sup>. However, data on its reproductive characteristics are still limited. The evaluation of reproductive performance in other fish species has had beneficial impacts in the development of freshwater aquaculture in

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Asia<sup>20,21,22,23,24</sup>. In contrast, there are still gaps in- knowledge of giant gourami sago strain broodfish regarding size at oocyte maturity, age of sexual maturity, sperm characteristics, egg hatchability, survival after eyed-egg stage, larva weaning and growth rate. These factors were identified as key challenges for successful sago strain giant gourami hatchery performance in the future. Therefore, the present study was conducted to evaluate the reproductive characteristics in <del>giant gourami</del>-sago strain gourami to provide basic knowledge for hatchery development for the future.

# Methods

#### Ethical considerations

There are no required permits from the government of the Republic of Indonesia to evaluate reproductive characteristics in sago strain gourami broodfish (*Osphronemus goramy*) as a candidate for future aquaculture. The study was founded by Research and Community Service Universitas Bung Hatta under a competitive grants scheme called the research of Professor in 2021 (Contract number: 06.02.1.46.03.2021). This grant included ethical approval and permits to collect fish specimens, rear and spawn sago strain gourami in the Aquaculture Laboratory Faculty of Fisheries and Marine Science Universitas Bung Hatta facilities. There was no animal suffering involved in this study sago strain gourami broodfish were still in good condition when returned to the pond. Ethical approval was granted by the Ethics Commission for Research and Community service at Universitas Bung Hatta (023/LPPM/Hatta/I-2021).

## Rearing and selection of breeders

Juvenile fishes were selected about six years ago from <u>a</u> local hatchery in Luhak District, Lima Puluh Kota Regency, West Sumatra Province. The juvenile fish were kept in tanks and transported by truck to the Aquaculture Laboratory, Faculty of Fisheries and Marine Science University of Bung Hatta. A total of 200 individuals <u>sago strain gourami</u> juvenile fishes were reared to sexual maturity\_in a concrete 8.4 m<sup>3</sup> (6×2×0.7 m) freshwater pond. During the rearing to sexual maturity, the juvenile fish were given commercial fish feed pellets (378-2 with 30% crude protein content and 4% crude fat<mark>; PT</mark> Japfa Comfeed Indonesia Ltd).

After that, a total of 30 <u>mature</u> individuals<u>s' adult sex</u> broodfish <u>were</u> separated according <u>by-to</u> sex and reared in two concrete <u>18 m<sup>3</sup> (6×2×1.5 m)</u> freshwater ponds <u>18 m<sup>3</sup> (6×2×1.5 m)</u>. Sago strain of giant gourami-broodfish<u>The broodfish</u> were feed twice daily (09:00 AM and 16:00 PM),<u>-were-with</u> extruded <u>pellet</u>feed <u>pellets</u> containing <u>was</u>-39.50% crude protein and 12.21% fat with a predetermined <u>ratio quantity as much asof</u> <u>-3% of biomass fish</u> weight per <u>/</u>day. Besides that, it is

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also were given sente leaves (Alocasia macrorriza L.) sufficiently as much as 1% (wet weight) of fish weight per day were contained 2.85% protein and 0.47% fat (% wet weight). Each concrete freshwater pond hads a 50 mm of middle drain in the middleage, which is was covered with a net of 0.5 cm mesh size to prevent the fish gurami sage broodfish from escaping and predators from entering. WThe water was pumped from the borehole wells at a velocity rate of 5 literes per minute.

A total of 20 broodfish for sage strain of giant gourami has oocyte matured with mature oocytes wereas selected, consistinged of 10 females and 10 males. Prior to stockingspawned, of females and males broodfish were weighed using balance scales (OHAUS model CT 6000-USA with 0.1 g accuracy), and their lengthsbody length were was measured using a meter ruler with 1mm accuracy. Average weight and length of the 10 female broodfish were 2140±159 g and 39.70±1.77 cm, while<sub>7</sub> 10 of those of the male broodfish were 3060±135 g and 43.1±1.79 cm.

Reproductionve characterization parameters in sage strain of giant gourami broodfishgiant gourami sage strain broodfish were analyzed using the following formulaes:

- Condition factor (CF) = wet weight in gram/length<sup>3</sup> × 100
- •\_\_\_\_;•O\_vulated egg weight (g) = fish weight before spawning (g) fish weight after spawning (g)
- •\_\_\_\_;••Ova somatic index (%) = egg weight ovulation (g)/ fish weight before spawning (g) x 100

#### Female reproductive performance

Starting in August 2020 onwards, the broodfish were checked each-monthly for eggs and semen production. The broodfish were captured with a hand net and anesthetized by orally with-ingestion of Tricaine methanesulfonate (MS-222, ethyl 4-aminobenzoate methanesulfonate 98%, Sigma Aldrich Co, USA, MO; 50 mg L<sup>-1</sup>), based on the dosage used for *Hemibagrus wyckii*<sup>21</sup>. Oocyte maturation was assessed for each individual. The oocyte maturity in sage strain of giant gourami females was assessed from oocytes sampled by intraovarian biopsies using a flexible polyethylene catheter<sup>21</sup>. Egg diameter was measured using Labo microscope model L-711 using and software camera 3.

Natural spawning of broodfish is-was carried out using in 1.2-\_m<sup>3</sup> (2×1×0.6 m) concrete freshwater ponds with a male-female sex ratio of (1:1). Before the broodfish is-spawned, the ponds are-were drained, cleaned and all other species are removed. Then, palm fibers are were provided which are placed on top of a bamboo raft in the pond. Furthermore, tThe pond is was then filled with water

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and the female and male broodfish <u>are-were</u> released into the spawning pond. The male broodfish made a nest for 5 to 7 days, after which spawning <u>takes-took</u> place and the female broodfish la<u>iyeds</u> eggs. Spawning <u>ed</u> occurred in the afternoon (<u>ranged frombetween</u> 15.00 to 17.00 PM). Due to the presence of a very large oil globule, <u>rage strein of glant gourami giant gourami sage strain eggs</u> float. After the broodfish ha<u>ds</u> finished laying eggs, the eggs <u>are-were</u> kept by the female broodfish in to the nest. <u>Furthermore, Aa</u>fter the eggs <u>are were had been</u> kept by the broodfish <u>for</u> four hours in the nest, the eggs <u>are-were</u> collected and transferred to an incubation tray, which <u>is-was</u> placed in a ventricular hatching system. A total of 100 eggs <u>eacheach broodfish are incubated in the</u> incubation trays, Meanwhile, the broodfish were returned to their pond <u>after-once</u> spawned, and no mortality occurred.

Egg weight and diameter were measured for 25 eggs per female using <u>balances (SHIMADZU-model</u> AY 220 <u>scales</u> with 0.1 mg accuracy) and <u>a miscrocopemicroscope (Labo microscope</u>-model L-711) using software camera 3. A total of 25 eggs were randomly sampled <del>at</del> 16 h<u>ours</u> after spawn<u>inged</u> to determine the fertility rate (FR). The hatching rate (HR) was determined by counting all hatched fry at 48 h<u>ours</u> after spawned. Then, endogenous feeding period counted until the egg yolks run out (day) and embryo survival rate (<u>%</u>) counted to eyed-egg stage <u>i.e. day 12.-(%)</u>

#### Determination of sperm quality

To stimulate the spermiation process in male broodfish used LHRH-a (Ovaprim, manufactured for Syndel Laboratories Ltd, 2595 Mccullough Rd Nanaimo B.C 9VS 4m9 Canada) Hormonal stimulation for spermination efficiency of males of sago strain of giant gourami broodfishgiant gourami sago strain broodfish used LHRH a preparations (Ovaprim) wwith dosage 0.5 ml per kg of brooder. Semen samples were obtained from 10 broodfish in sago strain of giant gourami randomly selected from the farm. The male broodfish were first anesthetized with 50 mg L<sup>-1</sup> of MS-222<sup>25</sup>, and-then, fish weights (MaW) and total lengths (MaL) were measured. Special care was taken to avoid any contamination of semen with urine, feces, mucus and water. Semen samples were collected using plastic syringes in 3 mL aliquots, and then placed in an insulated ice-cooled container, transported to the laboratory and analyzed within 2-two hours.

The sperm evaluate-assessment included gross (visual) and microscopic examination as reviewed by <u>Rurangwa et al. (2004) and Cabrita et al. (2017)</u><sup>26,27</sup>. The gross examination was based on visual and physical observation of parameters like the semen volume, semen pH, sperm concentration, motility and duration motility. Semen volume by collecting the semen in a graduated cylinder and determining the level in millimeters. Semen pH was determined with a hand pH meter (HI8424

Hanna Instruments, USA). The mMicroscopic examination was carried out using the Olympus model

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CX40, with <u>magnification between</u> × 10 and × 25 <u>magnification</u> to determine <u>other</u>-parameters such as motility (MO)\_percentage and duration,\_<u>were determined</u> by observing water\_activated semen placed on a glass slid<u>ee under a microscope</u>. Motile sperm were observed and expressed as a percent of non-moving sperm. Motility duration (DMO) was determined as the period between movements of the sperm to cessation of any progressive using Neubauer's hemocytometer and calculated as the number of sperm ml<sup>-128</sup>. <u>Semen pH was determined with a hand pH meter (HI8424</u> <u>Hanna Instruments, USA)</u>.

#### Water **q**Quality

Water samples were collected in the spawning pond and incubation trays were collected to determine the alkalinity, hardness and pH. <u>The protocol for determining alkalinity by standard</u> <u>methodology is presented by</u><sup>29</sup>. The traditional procedure for alkalinity is to measure how much H+ is required to titrate a sample to the methyl orange endpoint (about pH 4.5). The pH at the titration endpoint corresponds approximately to the point where an amount of H+ has been added to react with all the OH<sup>-</sup>, <u>CO3 2-</u>, and HCO3<sup>-</sup> in the sample to produce CO2 and H2O. The milliequivalents of H<sup>+</sup> used in the titration multiplied by 50.04 mg CaCO3/ Meq is the alkalinity. <u>Hardness is</u> <u>determined by titrating a sample with 0.01 M ethylenediaminetetraacetic acid (EDTA) to form</u> <u>complexes with divalent cations</u>. <u>Alkalinity and hardness were measured according to standard</u> <u>procedure presudure<sup>29</sup>, pH values were determined with a pH meter (digital mini pH meter, -)-, 14pH,</u> IQ Scientific, Chemo-science Thailand Co., Ltd, Thailand). An oxygen meter (YSI model 52, Yellow Spring Instrument Co., Yellow Springs, OH, USA) was used *in situ*, <u>and Water-water</u> temperature <del>of</del> <u>in</u> the spawning pond and incubation trays <del>as were m</del>easured with a thermometer (Celsius scale).

# Statistical analysis

Results were given as the <u>means-mean values</u> (± SD). Simple linear regression analyses were performed using <u>SPSS</u> software (version 16.0 for Windows; SPSS Inc., Chicago, IL). The standard deviation of each parameter was determined. For linear regression analyses, <del>significant</del> correlations were considered <u>significant</u> at p<0.05, and <del>whereas significant</del> trends or tendencies were considered <u>significant</u> at p<0.05.

# Results

The <u>reproduction reproductive</u> characteri<u>stics</u>zation of female broodfish in <u>sage strain of giantsage</u> <u>giant</u> gourami is summarized in Table 1. Total number of eggs <u>per nest was estimated per nest</u> (absolute fecundity) varied from 2000 to 2650, while relative fecundity (total number of eggs per kg

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female brooder) varied were between 977 and 1071. The fertility rate ranged from 76 to 84%, and the hatching rate-success rate ranged from 72 to 80%. Endogenous feeding period ranged from 10 to 12 days, and embryo survival rate to eyed-egg stage varied were between 94.73 and 95%.

Reproduction <u>Reproductive</u> characterization <u>characteristics</u> for males broodfish and sperm samples are presented in Table 2. The average live weight of the males <u>is-was</u> 3340±275.68 g. <u>Sago strain of</u> <u>Sago ge</u>iant gourami males broodfish <u>is-were</u> found to be slightly bigger than females broodfish. Gonad weight ranged from 25 to 30 g, whereas gonad somatic index ranged from 0.83 to 0.93%.

The analyzed of the linear correlation ( $r^2$ ) between variables of reproduction characteristicszation parameters in sago strain of giant gourami females broodfish <u>results are</u> shown in Table 3. <u>In t</u>This study, the <u>reproduction reproductive characterization</u> parameters that <u>had showed</u> strong correlations with the absolute fecundity were female fish weight before spawning ( $r^2 = 0.921$ ) and female fish weight after spawning ( $r^2 = 0.864$ ). Similarly, <u>results revealed significant correlations</u> between egg diameter <u>with and the</u> hardened egg diameter ( $r^2 = 0.833$ ), <u>egg diameter and</u> <u>percentage in the diameter of the hardened egg ( $r_{st}^2 = 0.699$ ), while egg diameter and fertility rate</u> <u>moderately correlated (hardened egg diameter increased ( $r^2 = 0.699$ ) and fertility rate ( $r^2 = 0.568$ ). In contrast, the egg diameter was not strongly related to absolute fecundity ( $r^2 = 0.169$ ) and relative fecundity ( $r^2 = 0.096$ ). On the other hand, the survival rate of larva<u>e</u>! (10 days)-is also had strong correlations with the hatching rate ( $r^2 = 0.998$ ) and endogenous feeding period ( $r^2 = 0.757$ ).</u>

The analyzed of the IL inear correlation analysis results ( $r^2$ ) between variables of reproduction characterization parameters in sago strain of giant gourami males broodfish are shown in Table 4. The reproductive characterization parameters that had a strong correlation with gonad weight were somatic index of gonads ( $r^2 = 0.836$ ), while semen volume ( $r^2 = 0.521$ ), semen pH ( $r^2 = 0.521$ ) and sperm concentration ( $r^2 = 0.506$ ) were moderately correlated with gonad weight. WhereasIn contrast, the gonadal weight negatively related correlated to with sperm motility ( $r^2 = 0.017$ ) and duration of motility ( $r^2 = 0.275$ ). Besides thatIn addition, the sperm concentration is was also had moderately correlated. with the sperm motility ( $r^2 = 0.556$ ) and duration of motility ( $r^2 = 0.502$ ).

The physico-chemical water quality parameters in the spawning ponds and- incubation trays for embryo development in sage strain of giant gourami-were as follows: water alkalinity ranged from 50.5 mg L<sup>-1</sup> to 52.5 mg L<sup>-1</sup>, hardness varied from 65.5 mg L<sup>-1</sup> to 67.5 mg L<sup>-1</sup>, pH ranged from 6.4 to 6.6, oxygen ranged from 6.1 mg L<sup>-1</sup> to 6.7 mg L<sup>-1</sup>, and- temperature varied from 28-°C to 30-°C.

Table 1. Reproduction characterization in sago strain of giant gourami females broodfish (Mean ±SD)

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	Variables	Range (Min-Max)		
Fish length (cm)	39.70±1.77	38-43	_	
ish weight before spawning (g)	2140±159.78	1958-2500		<b>Commented [CM25]:</b> This variable is missing from traw data; please clarify.
Fish weight after spawning (g)	2108±157.64	1930-2465		Commented [CM26]: This variable is missing from t
Condition factor	3.30±0.42	2.54-3.86		raw data; please clarify.
gg weight ovulated (g)	32.80±2.86	28-38		
Dva somatic index (%)	1.55±0.07	1.43-1.65		
Absolute fecundity (egg/fish)	2205±201	2000-2650		
Relative fecundity (egg/kg body weight)	1029±36	977-1071		
gg diameter (mm)	2.42±0.05	2.32-2.46		
lardened egg diameter (mm)	3.42±0.02	3.40-3.45		
gg diameter increase (%)	29.63±1.43	27.8-32.14		
gg weight (mg)	10.33±1.09	9.02-12.20		
Hardened egg weight (mg)	13.36±1.27	11.74-15.20		
Hardened egg weight increase (%)	22.69±2.24	19.74-24.81		
Fertility rate (%)	81.60±3.37	76-84		
Hatching rate (%)	76.40±2.27	72-80		
ndogenous feeding period (day)	11.2±0.63	10-12		<b>Commented [CM27]:</b> This variable is missing from the second seco
Embryo survival rate to eyed-egg stage (%)	94.76±0.42	94.73-95		raw data; please clarify. Commented [CM28]: This variable is missing from raw data; please clarify.

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# Table 2. Reproduction characterization in sago strain of giant gourami males broodfish (Mean $\pm$

SD)

	Variables	Range (Min-Max)	
Fish weight (g)	3060±134.99	2800-3200	
Fish length (cm)	43.1±1.79	40- 5	
Condition factor	3.74±0.43	3.08-4.38	

Gonads weight (g)	27.5±1.72	25-30
Gonadosomatic index (%)	0.90±0.03	0.83-0.94
Semen volume (mL per kg body weight)	0.60±0.12	0.4-0.7
Semen pH	8.18±0.15	7.9-8.4
Sperm concentration (10 <sup>9</sup> /mL)	1.44±0.14	1.2-1.6
Motility (%)	70.04±2.27	68-75
Duration of motility (sec)	50.2±7.25	43-61

	FEL	FWBS	FWAS	CF	OEW	OVI	AF	RF	EW	HEW	EWI	ED	HED	HDI	FR	HR	EFP
FEL																	
FWBS	<u>0.720</u>																
FWAS	<u>0.717</u>	<u>0.999</u>															
CF	<u>0.757</u>	<u>0.575</u>	<u>0.574</u>														
OEW	<u>0.539</u>	<u>0.565</u>	<u>0.553</u>	0.365													
OVI	0.281	0.191	0.012	0.000	<u>0.774</u>												
AF	0.255	<u>0.921</u>	<u>0.864</u>	<u>0.637</u>	0.387	0.072											
RF	0.012	0.207	0.063	0.011	0.065	0.321	<u>0.524</u>										
EW	<u>0.894</u>	<u>0.659</u>	<u>0.655</u>	<u>0.677</u>	<u>0.552</u>	0.246	<u>0.567</u>	0.041									
HEW	<u>0.841</u>	<u>0.631</u>	<u>0.626</u>	<u>0.514</u>	<u>0.582</u>	0.295	0.468	0.004	<u>0.924</u>								
EWI	0.165	0.109	0.109	0.354	0.033	0.000	0.010	0.002	<u>0.924</u>	0.041							
ED	0.164	0.132	0.338	0.131	0.378	<u>0.688</u>	0.169	0.096	0.030	0.263	0.000						
HED	0.064	0.029	0.237	0.126	0.025	0.207	0.022	0.064	0.468	0.184	0.030	<u>0.833</u>					
HDI	0.266	0.293	0.342	0.209	0.103	0.085	0.373	0.195	0.318	0.298	0.006	<u>0.699</u>	0.294				
FR	0.026	0.229	0.020	0.000	0.135	0.264	0.000	0.004	0.067	0.004	0.160	<u>0.568</u>	0.064	0.054			

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HR	0.035	0.020	0.060	0.046	0.000	0.009	0.003	0.009	0.226	0.108	0.364	0.018	0.001	0.143	<u>0.703</u>		
EFP	0.113	0.186	0.190	0.094	0.006	0.098	0.231	0.103	<u>0.747</u>	<u>0.806</u>	0.147	0.317	0.001	0.116	0.015	0.013	
SR	0.070	0.032	0.034	0.027	0.007	0.003	0.112	0.024	0.194	0.019	0.005	0.033	0.000	0.324	0.063	<u>0.998</u>	<u>0.757</u>

Statistically important at  $r^2 > 0.500$  (underlined)

FeL: female fish length (cm); FWBS: female fish weight before spawning (g); FWAS: female fish weight after spawning (g); CF: condition factor; OEW: egg weight ovulation (g), OVI: Ova somatic index (%), AF: absolute fecundity (eggs), RF: relative fecundity (eggs), EW: egg weight (mg), HEW: Hardened egg weight (mg), EWI: eggs weight increase (%), ED: egg diameter (mm), HED: hardened egg diameter (mm), HDI: hardened egg diameter increase (%), FR: fertility rate (%), HR: hatching rate (%), EFP: endogenous feeding period (days), SR: survival rate (10 days).

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Table 4. Correlations of variables (r<sup>2</sup>) in sago strain of giant gourami males broodfish

	Mal	N4-14/	<b>CГ</b>	CIN	CI	<u>C) /</u>		50	140
	MaL	MaW	CF	GW	GI	SV	рН	SC	MO
MaL	-								
MaW	<u>0.714</u>								
CF	<u>0.807</u>	0.347							
GW	0.399	<u>0.550</u>	0.187						
GI	0.003	0.042	0.071	<u>0.836</u>					
SV	0.025	<u>0.576</u>	0.042	<u>0.521</u>	0.000				
рН	<u>0.516</u>	<u>0.772</u>	0.353	<u>0.521</u>	0.127	0.296			
SC	0.186	<u>0.661</u>	0.131	<u>0.506</u>	0.068	0.425	<u>0.645</u>		
MO	0.068	0.453	0.061	0.017	0.130	0.393	0.280	<u>0.556</u>	
DMO	0.159	0.322	0.083	0.275	0.012	0.082	0.430	<u>0.502</u>	<u>0.519</u>

Statistically important at  $r^2 > 0.500$  (underlined)

MaW: male fish weight (g), MaL: male fish length (g), CF: condition factor; GW: gonadal weight (g); GI: gonad somatic index (%), SV: semen volume (ml), SC: sperm concentration (10<sup>9</sup>/mL), MO: motility (%), DMO: duration of motility (sec).

Table 5. Summary of the fecundity, gonadal somatic index, egg diameter and hatching rate of giant	
gourami.	

Species	Strain	Relative	GSI (%)	Eggs	Hatching	Reference
		fecundity		diameter	rate (%)	
		(egg/kg fish)		(mm)		
Osphronemus	Sago	1037±90	1.91±0.35	2.42±0.05	76.40±6.33	This study
goramy						
Osphronemus	Bastar	2423±348	2.78±1.16	2.2 ± 0.2	96.36± 2.30	16
goramy						
						8
Osphronemus	Galunggung	4011±287	4.15±0.63	2.5±0.05	89.3±1.30	0
goramy						

Osphronemus goramy	-	5508±1547	2.32±0.50	2.18±0.19	61.60±0.0	30
Osphronemus	Tambago	2.896±185	3.16±0.11	2.47±0.03	91.06±4.06	31
goramy						

## Discussion

In our study, body weight in <u>female</u> sago <u>strain of giant</u> gourami <u>female</u> broodfish before spawning ranged from 1958 to 2500 g per fish and ova somatic index ranged from 1.43 to 1.65%. Body weight in <u>female</u> sago <u>strain of</u> giant gourami <u>female</u> broodfish <u>was is the</u> smaller than as that of giant gourami belonging to the galunggung strain, <u>which</u> range<u>de</u> from 2500 to 3500 g<sup>§</sup>. Conversely, ova somatic index of galunggung strain are found to be slightly bigger than <u>with that of</u> sago <u>strain of</u> giant gourami, <u>which</u>-ranged from 3.7 to 4.6%<sup>8</sup>. The differences <u>of in</u> reproductive<del>on</del> <del>characterization</del>-<u>characteristics</u> in broodfish can be <u>affected explained</u> by strains, brood size, <u>age of</u> <u>broodfishused</u>, previous spawning history and the production setting<sup>23</sup>.

Absolute fecundity (AFs) in sago strain of giant gourami ranged between from 2000 to 2650 eggs fish<sup>-1</sup> and <u>relative fecundity (RF)</u>s were ranged from 977 to 1071 eggs kg<sup>-1</sup>. Egg produced in kg fish<sup>-1</sup> (RF) is thought to be more informative than absolute fecundity. Relative fecundity (RF) in sago strain of giant gourami were smaller compared to those in galunggung strain, palapah strain and blusafir strain<sup>8,31,16</sup>. On the other hand, the difference of relative fecundity can also be related to differences in broodfish size and age used<sup>23</sup>. Environmental factors such as rainfall is-also influenced the number of eggs per spawn in giant gourami brood, while the water temperature negatively related to number of eggs per spawn<sup>15</sup>. Furthermore, egg diameter in sago strain of giant gourami is found to be almost the same than others strain of giant gourami (Table 5). In this study,-EDS egg diameters obtained average was 2.42±0.05 mm, consistent with those reported by another researchers, for example 2.18±0.19 mm for the giant gourami<sup>30</sup>, 2.40±0.05 mm for blusafir strain<sup>16</sup> and 2.5±0.05 mm for galunggung strain<sup>8</sup>. Apparently, tThe differences the relative fecundityRF, ova somatic index, egg diameter and hatching rate of giant gourami can be influenced by differences in the strains. Furthermore, the egg diameter has been influenced by the dietary protein level<sup>32,33,34,35</sup>, age of broodfish<sup>36</sup>, and spawning season<sup>37,38</sup>. In our study, egg diameter was shown to be positively correlated with egg weight, hardened egg weight, and eggs weight increased. Egg weight of rainbow trout also increased after the hardening process and is a positively correlated with the viability of

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# eggs<sup>39</sup>. Other egg quality metrics, such as hatching rate, and survival to first feeding, has been correlated with good egg quality<sup>21</sup>.

In this study, the hatching rate of embryo (HRs) in sago strain of giant gourami were smaller than those of other strains of giant gouramy<sup>8,31,16</sup>. This condition might be affected by the egg and sperm quality in sago strain of giant gourami broodfishgiant gourami sago strain broodfish. In the present study, whether eggs and sperm quality of sago giant gourami breeders are affected by feed nutrition type for sago strain of giant gourami brooder wasis poorly understood. Meanwhile, the bB roodfish sex ratio has nothad no influence d-on egg quality<sup>8</sup>. The reproduction reproductive characterization parameters that had strong correlations were strongly correlated with the hatching rate were fertility rate ( $r^2$ =0.703) and survival rate (10 days) ( $r^2$ = 0.998). According to Sink *et al.*<sup>40</sup> that biochemical composition of egg strong correlated with egg quality of broodfish. According to Sink *et al.*<sup>40</sup> that al. (2010)<sup>40</sup> that indicator of egg quality the strong correlation with biochemical composition of eggs. In this study, we did not evaluate the biochemical composition of egg, because -due to-the relationshiped between egg quality and biochemical composition is difficult to interpret<sup>41</sup>.

The keys of regulator <u>hormoness</u> of fish reproductioned were are gonadotropins (GTHs), folliclestimulating hormone (FSH), luteinizing hormone (LH), and sex steroids<sup>42,43</sup>. Besides that<u>In addition</u>, oocyte develop<u>mented</u> and matur<u>ationed are</u> also regulated by locally acting <u>paracrine and</u> autocrine <u>signalling</u><sup>14,45</sup>. However, there is not information about the effects of such kinds of factors on the oocyte develop<u>ment</u> in sago <u>strain of</u> giant gourami. <u>Conversely, the extruded feed enriched</u> with vitamin E (d- $\alpha$ -tocopherol) as much as 137.8 mg, 238.05 mg, 338.72 mg and 439.39 mg per kg of feed ingredients had an effect on markers of brood reproduction function. gouramy, such as <u>sexual maturity time</u>, ovum somatic index, relative fecundity and egg diameter31.<u>Nevertheless</u>, extrusion feed enriched with the vitamin E (d- $\alpha$ -tocopherol) as much as 137.8, 238.05, 338.72 and 439.39 mg per kg feed ingredients affected to markers of reproductive functions of giant gourami brooder (e.g., time sexual maturity, ova somatic index, relative fecundity and egg diameter]<sup>31</sup>.

Various efforts have been made by scientists to increased reproductive performance of female broodfish, such as to increasing e about dietary protein level for *Xiphophorus helleri*<sup>33</sup>, *Channa marulius*<sup>46</sup> and *Ictalurus punctatus*<sup>40</sup>. Additionally, implantation of 17ß-estradiol has also managed to improved the reproductive performance in *Hemibagrus nemurus*<sup>47</sup>. <u>Currently In this context</u>, whether the increase <u>in</u> protein level of feed and use <u>of</u> hormon<u>esal</u> can increase <u>the</u> reproduc<u>tiveed</u> potential in sago <u>strain of</u> giant gourami <u>are is</u> poorly understood. Therefore, we recommend the use <u>experiment use</u> of feed protein levels and hormonal dosage <u>to increase the reproductive potential it</u>

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is very important toin sago strain gourami broodfish is very important to be applied applied for sago strain of giant gourami broodfishgiant gourami sago strain broodfish in the future.

Average SVs semen volume was found in sago strain of giant gourami are were lower (0.4 to 0.6 ml) than those of *Hemibagrus wyckii* (0.60 to 1.20 ml)<sup>21</sup>, but higher than those of *Pterygoplichthys gibbiceps*<sup>48</sup>. It appears that the semen volume depend on fish species<sup>21,49,50</sup>. So mMany factors influenced onsperm quality and quantity of sperm such as genetic, physiological, spawning season and environmental factors.<sup>26,49,51,52</sup>. On the other hand, improvements nutrition feed of broodfish can be and proofer feeding greatly improved gamete quality and larval semen production volume<sup>46</sup>. Commercial honey combined with 10% Dimethyl Sulfoxide (DMSO) it is also had increased motility rate of sperm<sup>53</sup>. While, synthetic hormones such as salmon gonadotropin releasing hormone analogue and domperidone (GnRH + Domperidone) effective on sperm quality<sup>54,55</sup>. Nevertheless, for sperm fertility competition occurs in the aquatic environment, and that had strong correlation and was strongly correlated with the water quality in ponds. In this study, the water quality parameters in spawning ponds of sago strain gourami broodfish in terms of alkalinity (50.5 to 52.5 mg L<sup>-1</sup>), and hardness (65.5 to 67.5 mg L<sup>-1</sup>), pH (6.4 to 6.6) and water temperature (28 to 30 °C). All of these water quality parameters have been able to support the ability of sperm to fertilize an egg.

The all water quality parameters can be supported the ability for sperm to fertilized an egg.

The sperm motility in sago strain of giant gourami ranged from 68 to 75%, and duration of motility ranged from 43 to 61 sec. Theseis results that are consistent with Genypterus blacodes and Esox lucius<sup>56,50</sup>. The Seperm motility includesed the percentage of motile sperm, straight line velocity, curvilinear velocity, average path velocity and linearity<sup>50</sup>. In this study, we did not investigate of those parameters. MeanwhileIn addition, the percentage of motileity sperm is influenced by addition of extenders and cryoprotectants<sup>57,58,59</sup>. However, sperm motility on from fresh semen was slightly better greater compared to cryopreserved vation semen on from the Esox lucius<sup>50</sup>. The fertility rate of eggs ranged from 76 and 84%; neverthelesshowever, it is does not detected any no significant<del>tly</del> correlation was detecteds between fertility rate and sperm parameters, such as semen volume, semen pH, motility and duration motility. Conversely, the sperm concentration had was moderately correlated ion with the sperm motility and duration of motility. The parameters commonly measured to assess sperm quality in brood were volume, density and motility (such as the percentage of motile sperm, straight line velocity curvilinear velocity, average path velocity, linearity and amplitude of lateral head displacement), includinged fertilizing capacity 49,50,52,60. In **∓**<u>t</u>his study, we did not investigate the ionic composition of the semen, but this phenomenon could be related to the ionic composition of semen which might has a significant influence on sperm motility and duration of motility.

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# Conclusions

This research analyzed the reproduction reproductive characterization characteristics in the sage strain of giant gourami broodfishgiant gourami sago strain broodfish reared in concrete freshwater ponds, in the Aquaculture Laboratory Faculty of Fisheries and Marine Science, Universitas Bung Hatta. Relative fecundities of the Sago strain of giant gourami broodfishgiant gourami sago strain broodfish had the relative fecundities ranged from 977 to 1071 eggs, and egg diameter between ranged from 2.32 to 2.46 mm. The semen-Semen volume ranged from 0.4 to 0.7 ml per kg body weight and sperm motility were-was comprised between 68 to 75%. There was a A strong linear relationship<u>was observed</u> between absolute fecundity with and the female fish weight before and after spawning. Similarly, the A similar strong correlated relationship was observed between absolute fecunditysurvival rate (10 days) with hatching rate. Sperm parameters which had moderately correlated was between sperm concentration with sperm motility and sperm duration of motility, and male sperm concentration and sperm motility for the reared sago strain. Keys to increasing reproduction performance in sago-strain of giant gourami dependes upon on broodfish weight-size of broodfish, relative fecundity and hatching rates. Despite initialdata on reproductive hascharacteristic has been success for spawningobtained, there is still an are-observed limitations of quality seed supply for aquaculture attributed to knowledge gaps in larval weaning, grow-up feeding technologies. Therefore, for the success successful practiced practices in hatcheryhatcheries, where further research is recommended for-to determine a proper feed formulation and the development

of appropriate aquaculture systems.

#### **Data availability**

#### Underlying data

RepositoryFigshare: Title. Reproduction characterization of the gurami sago (*Osphronemus goramy* Lacepède, 1801): basic knowledge for a hatchery development strategy for the future. https://doi.org/10.6084/m9.figshare.14661189.v1 [61].

DOI: 10.6084/m9.figshare.14661189

This project contains the following underlying data:

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- Table 1. Raw data of fish length, weight, absolute fecundity and relative fecundity of gurami sago broodfish
- Table 2. Raw data of egg diameter (mm) in sago strain of giant gourami broodfish
- Table 3. Raw data of hardened egg diameter (mm) in sago strain of giant gourami broodfish
- Table 4. Raw data of egg diameter increase (%) in sago strain of giant gourami broodfish
- Table 5. Raw data of egg weight (mg) in sago strain of giant gourami broodfish
- Table 6. Raw data of hardened egg weight (mg) in sago strain of giant gourami broodfish
- Table 7. The data of egg weight increase (%) in sago strain of giant gourami broodfish
- Table 8. The data of fertilization rate (%) in sago strain of giant gourami broodfish
- Table 9. The data of hatching rate (%) in sago strain of giant gourami broodfish
- Table 10. Male size, gonadal weight and semen in sago strain of giant gourami broodfish
- Table 11. Sperm concentration (10<sup>9</sup>/mL) in sago strain of giant gourami broodfish
- Table 12. Sperm Motility (%) in sago strain of giant gourami broodfish
- Table 13. Duration motility (sec) in sago strain of giant gourami broodfish

Data are available under the terms of the Creative Commons Attribution 4.0 International license (CC-BY 4.0). License: CC-BY 4.0

# **Competing interests**

No competing interests were disclosed.

# Grant information

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