# A Novel Low Temperature Synthesis of Sodium Silicate (Water Glass) from Silica-Rich Wastes using Hydrothermal Method

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## Abstract

This present work aims at synthesizing sodium silicate solutions from selected industrial wastes (waste glass and rice husks) using hydrothermal method. The as-received waste glasses of different mix colors were initially washed, crushed and wet milled for several hours then sieved through 75µm mesh. The sieved recycled glasses were then oven dried at 110°C for 72 h to obtain a powdered glass (SLSG). Rice husks were also washed to remove dirt adherence and dried. The dried rice husks were thermally treated in a muffle furnace at 800°C for 1 h then left to cool to obtain whitish rice husk ash (RHA). Both SLSG and RHA were then reacted with NaOH solution at varying concentration of 3M, 4M and 5M respectively in a 250 ml Erlenmeyer flask. The mixtures were heated with continuous stirring using a magnetic stirrer at temperatures of 150°C, 170°C and 200°C respectively for a reaction period of 4 h. Analyses were carried out on the synthesized sodium silicate solutions. The results obtained from the analyses showed sodium silicate synthesized from RHA possessed high viscosity while the SiO<sub>2</sub>/Na<sub>2</sub>O ratio showed they are commercial sodium disilicates which indicates the products are all sodium silicate solutions. **Keywords**: Sodium silicate; industrial wastes; hydrothermal; viscosity; chemical composition.

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### 1. Introduction

Sodium silicate solutions (Na<sub>2</sub>SiO<sub>3</sub>) which are commercially referred to as water glass are liquids like dissolved glass containing complex mixtures of anions and polymer silicate particles of silica/sodium oxide (SiO<sub>2</sub> : Na<sub>2</sub>O) molar ratio > 2 and possess some water like properties [1]. Sodium silicate is used for broad range of industrial applications such as synthesis of gas adsorbents (silica xerogel) to trap carbon (IV) oxide [2], zeolites and metallic organic framework materials [3], aerogels [4], ingredient in cement production [5]. Other uses of sodium silicate include applications in detergents, corrosion inhibitor, sealants, deffloculants, binders, paper and pulp etc.

Industrially, production of sodium silicate involves fusion of mixtures of sand (SiO<sub>2</sub>) and soda ash (Na<sub>2</sub>CO<sub>3</sub>) at elevated temperature above 1200°C resulting into formation of glass on cooling; the glass formed is then dissolved in water to produce soluble sodium silicate which is commercially known as water glass [6, 7]. However, this process of synthesizing sodium silicate solutions is considered to be quite costly due to high energy consumption required to attain high temperature and entails negative environmental effects resulting from emission of pollutants such as dust particles, sulfuric oxide, nitrogen and carbon monoxide from decarbonation process. As a result of this, a new and cheaper approach was developed in regard to the chemistry of silica (green chemistry) which involves synthesis in a way that is sustainable, safe, and non-polluting and which consumes minimum amount of materials and energy while less or no waste material is produced. This process involves reaction of silica rich material with aqueous solution of sodium hydroxide (NaOH) in an open system under atmospheric pressure using lower temperature as adopted by several works and patents [8 - 12]. In recent times, crucial economic and environmental considerations have geared industries and researchers to develop and improve technologies targeted at drastically minimizing industrial wastes. In this regard, several efforts have been devoted on the utilization of wastes which are known to be highly rich in silica emanating from urban and industrial sectors as alternative to natural silica. In Nigeria, more than 32 million tons of solid wastes are annually produced from various municipal and industrial processes, of which waste glasses and agro wastes account for 70 percent. The usual practice in this country as a developing nation is to primarily treat these wastes in incineration plants and which has raised a lot of environmental concerns. Though waste glasses and rice husks are non-toxic but their accumulations are subject of concern presently in the country. Hence, efforts have now been attached toward recycling and valorization of wastes and by-products in making entirely new and useful products which in turn serve as a means of environmental protection [13]. Recently, several authors have investigated and reported the use of silica rich waste materials as silica source in the synthesis of sodium silicate [7 - 9, 14]. However, these investigations concentrated on agro-waste materials while little work on the use of waste glasses [13, 15] utilized single colored waste glasses without varying parameters such as concentration and temperature. Therefore, given consideration to the cost incurred and time of sorting these waste glasses into their different colors, this research adopted the use of mix colored waste container soda-lime-silica glasses through blending of three major colors (green, brown and white). This will not only reduce cost of sorting and beneficiation but also ensure more glasses are recycled within short period.

This present work therefore aimed to synthesize soluble sodium silicate using SLSG (soda-lime-silica glass) and RHA (rice husk ash) as silica sources and sodium hydroxide (NaOH) as soda precursor instead of conventional soda ash (Na<sub>2</sub>CO<sub>3</sub>) at different temperatures and concentration of NaOH, then make a comparative study between the sodium silicate obtained from the two silica sources (SLSG and RHA) while chemical composition and ideal physical properties such as viscosity, <sub>P</sub>H and specific gravity were investigated.

## 2. Materials and methods

## 2.1. Materials

The starting materials used in this work are waste container soda-lime-silica glasses (SLSG) of different colors (green, brown and white), rice husks (RH) and sodium hydroxide pellets (purity 98%). The SLSG and RH serve as silica sources while sodium hydroxide pellets is used as precursor to sodium oxide (Na<sub>2</sub>O) instead of soda ash. The sodium hydroxide was purchased from a reliable chemical vendor and used in its as-received state. The SLSG of different colors were sourced from municipal dumpsite while rice husks were obtained as agro wastes from rice milling plant. The as-received SLSG (green, brown and white) were initially washed thoroughly and allowed to dry properly prior to crushing into smaller particle sizes using a laboratory jaw crusher. After crushing, the glasses were then poured with addition of water inside a porcelain-lined ball mill using porcelain balls as milling media; this is to prevent contamination. The milling was done for 48 h to obtain glass slurries which were later sieved through 75µm mesh. Wet milling was adopted to prevent excessive attrition. The wet-milled recycled glasses were then dried in an electric oven for 72 h to obtain powdered recycled SLSG. It was possible to blend all the colored glasses together because all container glasses are of similar composition with only variation from the coloring agents in the colored glasses. However, this coloring agents in the colored glasses cannot altered the synthesis of sodium silicate. This practice also eliminates the bottlenecks involved in color sorting thus reducing cost and saving time. For the rice husks, the as-received rice husks were initially washed to remove adhered dirt followed by air drying for 72h. The dried husks were then stacked inside incinerator for combustion into carbonized rice husk ash. The carbonized rice husk ash was further treated thermally to a temperature of 800°C inside a muffle furnace for a holding period of 1h at a heating rate of 10°C/min to obtain a pure white rice husk ash (RHA) which was then sieved through 75µm sieve to obtain a fine powder RHA. Both the powdered SLSG and RHA were then characterized following standard procedures. Chemical composition of the SLSG and RHA was examined using high performance Energy Dispersive X-ray Fluorescence (EDXRF) XSUPREME 8000 to quantify the concentrations of SiO<sub>2</sub> and other oxides present. Phase identifications were determined by X-ray diffractometer using BRUKER AXS with D8 Advanced diffractometer Cu Ka radiation XRD in the range of 2 tetha angle from 5 to 80 scanning range. Identification of molecular bonding was evaluated using Fourier-transform infrared spectroscopy (spectrum 100 FT-IR Spectrometer, Perkin Elmer) while the specific surface area of both recycled wastes were determined using Brunauer - Emmett - Teller (BET) analyzer. Figures 1 (a - d) showed the as received wastes and recycled wastes respectively.

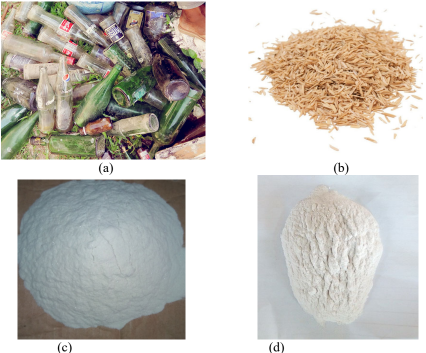


Figure 1: Representative pictures of (a - b) recycled wastes, (c - d) processed powders

## 2.2. Synthesis of sodium silicate (hydrothermal method)

The hydrothermal method used in this work is an open system following the adopted method of Nittaya and Apinon [16] under atmospheric pressure (14.70 psi). The system consists on the use of glass reaction vessel, heating source, temperature measuring device and a reflux condenser, to ensure constant volume of the reaction mixture. In this work, appropriate amount (10 grams) of recycled SLSG and RHA were at separate time mixed with 80 ml of 3.0, 4.0 and 5.0 M sodium hydroxide solutions respectively in a 250 ml Erlenmeyer flask. The flask containing the mixture was placed on a hot plate connected with a reflux condenser and continuously stirred with the aid of magnetic stirrer at 40 rpm under varying reaction temperature of 150°C, 170°C and 200°C for a period of 4h. Constant stirring was ensured to dissolve the silica and produce sodium silicate solutions. The solutions were then filtered using a Whatman No. 41 filter paper to obtain sodium silicate solutions while the residues were washed with 50 ml boiling water.

#### 2.3. Characterization

Physiochemical analyses such as chemical analysis, viscosity, pH, specific gravity were investigated to assess the synthesized sodium silicate solutions. The pH of the synthesized sodium silicate solutions were analyzed using a pH meter (Oakton pH 700 Benchtop Meter) following standard procedures while viscosity of the sodium silicate solutions were also analyzed with the use of viscometer (Brokfield DV III Programmable Rheometer) at 25°C. The chemical analysis of the sodium silicate solutions was carried out to determine the percentage of sodium oxide (Na<sub>2</sub>O) and silica (SiO<sub>2</sub>) which are two major components of typical sodium silicate present in the synthesized sodium silicate solutions. The SiO<sub>2</sub> was characterized using the American Public Health Association (APHA) 4500-SiO<sub>2</sub> C. Molybdosilicate Method [17]. Standard solutions of silica of variable concentrations; 5-25 ppm were prepared from silica stock solution. UV/VIS Spectrometer scan (SpectroLab 753S UV/VIS Spectrometer) of the standards were run and the absorbance measurements at 410 nm were used to prepare the calibration curve. The prepared samples were diluted to 2.5  $\mu$ l in 50 ml distilled water (2.5 mg in 50 g solution: dilution factor; 5 x 10<sup>-5</sup>). The diluted solutions of the samples were mixed with ammonium molybdate. The pH of the resulting mixture was adjusted to 1.2 by adding 1 ml of 1 + 1 hydrochloric acid. The mixture was allowed to stand for 10 min then 2.0 ml of oxalic acid was added. UV-VIS absorbance at wavelength of 410 nm was taken and the concentration of the sodium silicate was determined. The corresponding concentration of the silicate, in ppm, was determined using the calibration curve. The actual concentration of the silicate, in wt%, was determined using the equation: Actual conc. (wt) = (conc. (ppm) x dilution factor)/  $(1 \times 10^{-4})$ .

The Na<sub>2</sub>O was characterized using Flame Photometer analysis (SpectroLab Flame Photometer SL-160). Firstly, variable concentrations of standard solutions of Na<sub>2</sub>O were prepared; 5-25 ppm from Na<sub>2</sub>O stock solution. The standard solutions were introduced into the flame photometer for inbuilt calibration. The sodium silicate solution was diluted to 2.5  $\mu$ l in 100 ml distilled water (2.5 mg in 100 g solution: dilution factor; 2.5 x 10<sup>-5</sup>) and introduced into the flame photometer.

#### 3. Results and Discussion

The chemical compositions of the recycled SLSG and RHA carried out by EDXRF in the form of stable oxides are presented in Table 1. It can be observed that both recycled wastes show similar chemical composition with major component being silica (SiO<sub>2</sub>). However, the highest silica content of 80 wt. % is obtained for RHA while SLSG has 69.4 wt. % SiO<sub>2</sub>. The chemical composition of RHA especially the amount of SiO<sub>2</sub> is a function of combustion process, temperature, harvest year of rice husks etc. as stated by [18] while that of SLSG varies from one factory to the other. However, the SiO<sub>2</sub> content obtained indicates that both wastes have high silica content to be used as silica sources for production of sodium silicate.

Figures 3 (a – b) indicate the results of the phase constituents of the SLSG and RHA by XRD. It is observed that SLSG showed a single broad band peak between  $15^{\circ} - 30^{\circ}$  as shown in Figure 5(a) which indicates that SLSG is amorphous [19]. However, in Figure 5(b), the diffractogram of RHA indicates peaks of cristobalite and quartz while tridymite also exist alongside other oxides. The initial broad band peaks observed between  $5^{\circ} - 20^{\circ}$  in the diffractogram indicate that RHA exhibited slight amorphous nature [19]. However, amorphous silica has been identified to be most soluble form of silica [20 – 21] which is very relevant to the synthesis of sodium silicate for easy dissolution of the silica constituent.

The results of the molecular bonding existing in the SLSG and RHA are presented in Figures 4 (a) and (b) respectively. The IR spectra of the SLSG and RHA were recorded by FTIR at wavenumber ranging from 500 to 4000 cm<sup>-1</sup>. The FTIR spectra were similar with two predominant transmittance peaks at certain wavenumber in the spectra. The transmittance peaks of RHA were observed at 1069.7 cm<sup>-1</sup> and 797.7 cm<sup>-1</sup> while the transmittance peaks of SLSG were recorded at 939.3 cm<sup>-1</sup> and 767.8 cm<sup>-1</sup>. However, additional weak peaks were detected for SLSG at 1420 cm<sup>-1</sup>, 2050 cm<sup>-1</sup> and 2139.5 cm<sup>-1</sup> respectively. The predominant intensity peaks at wavenumber range of 900 – 1100 cm<sup>-1</sup> that was observed in both RHA and SLSG might be attributed to Si – O – Si asymmetric bending and stretching vibration bonds indicating the presence of siloxane structural bonding as reported by [22 –

23]. This is in similarity with what was observed for synthetic silica reported by [24]. Weak peaks recorded at wavenumber range 700 - 800 cm<sup>-1</sup> in the spectra of both RHA and SLSG can be attributed to stretching of symmetric Si – O bond referred to as Silanol as reported by [22]. The additional weak peak observed in SLSG at 1420 cm<sup>-1</sup> and wavenumber range 2000 - 2500 cm<sup>-1</sup> might be due to absorption and disturbed silanol respectively.

Table 2 showed the results obtained from the BET analysis respectively. From the data obtained, it is observed that RHA has the highest BET total surface area of 22.3 m<sup>2</sup>/g while SLSG has 1.9 m<sup>2</sup>/g. The highest surface area observed for RHA indicates that a large portion of the RHA will react with NaOH resulting into better dissolution of silica resulting into quality sodium silicate solutions. However, SLSG has the highest micropore area of 1.3 m<sup>2</sup>/g.

The results of the pH measurements of the synthesized sodium silicate solutions from soda-lime glass and rice husk ash are shown in Table 3 and 4 respectively. The results of the pH measurements showed that the pH values of the synthesized sodium silicates for both SLSG and RHA increases with alkali (NaOH) concentration of 3M, 4M and 5M used which is actually expected. Also, in relative to synthesizing temperature ( $150^{\circ}$ C,  $170^{\circ}$ C and  $200^{\circ}$ C) there is just slight drop in pH values for RHA synthesized sodium silicate while slight increase in pH values for SLSG synthesized sodium silicates can be observed. However, there is no typical standard for pH range of sodium silicate solutions as it is a function of alkali concentration. But, considering the pH values obtained in this research for both SLSG and RHA synthesized sodium silicate solutions, it can be observed that the pH values fall within 13.0 - 14.0 which is an indication of high alkalinity. In regard to this high alkalinity, the sodium silicate solutions synthesized in this research can be recommended for industrial applications such as soaps/detergent, xerogel, silica extraction and as a clay deflocculant.

The results of the representative viscosity measurement are shown in Table 5. Three best samples were selected for sodium silicate synthesized from SLSG while three best samples were also selected for sodium silicate synthesized from RHA. The selection parameter was based majorly on the visual observation of the products in term of viscousness. From the viscosity measurements, it can be observed that selected samples possessed considerable viscosity though less than 330 Pa.s for commercial sodium silicate [25]. However, it can be observed that sodium silicate solutions synthesized from rice husk ash possessed high viscosities above 300 Pa.s falling within the range of commercially produced sodium silicate solution which might be attributed to the high surface area of reaction with NaOH resulting into better silica dissolution thus enhancing viscosity. However, the viscosities of samples obtained from SLSG are still considerable.

The results of the chemical analysis of the best samples selected are shown in Table 6. In this work,  $SiO_2/Na_2O$  ratio was also calculated from the results of the chemical analysis obtained to determine the reaction efficiency and to affirm the types of sodium silicate obtained. Based on literature, the type of sodium silicate solutions synthesized is a function of the  $SiO_2/Na_2O$  ratio obtained [26]. It can be observed that the amount of silica (SiO<sub>2</sub>) extracted from the wastes (SLSG and RHA) used as silica sources increased as the synthesized temperature increased from  $150^{\circ}C - 200^{\circ}C$  which indicates that silica dissolution from both SLSG and RHA depend majorly on the temperature. However, the amount of soda (Na<sub>2</sub>O) dissolved is a function of the concentration of the NaOH used as source of sodium oxide. It is observed from the SiO<sub>2</sub>/Na<sub>2</sub>O ratio that all the best samples selected fall within 1.89 - 2.45 and are therefore referred to commercially as sodium disilicate. In term of SiO<sub>2</sub>/Na<sub>2</sub>O ratio, samples having high SiO<sub>2</sub>/Na<sub>2</sub>O ratio is preferable for industrial synthesis of silica gel as small amount of acid will be required for formation of silica gel [26]. On the average, sodium silicate synthesized from RHA has the highest SiO<sub>2</sub> content due to their large surface area exposed for reaction.

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	SiO <sub>2</sub>	Al <sub>2</sub> O <sub>3</sub>	CaO	MgO	Na <sub>2</sub> O	K <sub>2</sub> O	P2O5	Fe <sub>2</sub> O <sub>3</sub>	$Cr_2O_3$	Mn <sub>2</sub> O <sub>3</sub>	TiO <sub>2</sub>
RHA	80.0	1.48	1.69	3.12	0.08	2.58	9.49	1.09	0.00	0.14	0.09
SLSG	69.4	5.03	15.03	0.55	7.22	0.65	0.09	0.71	0.13	0.02	0.65

Table 1. Chemical composition of the RHAs and WSLSG by EDXRF

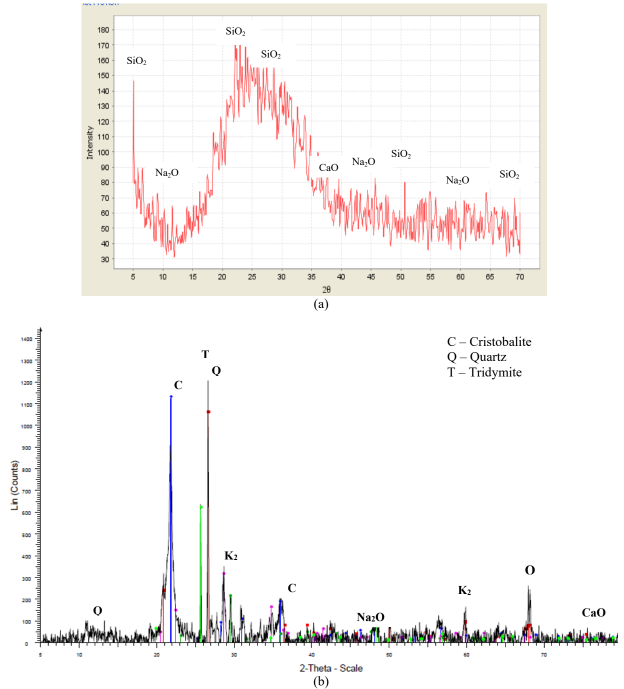


Figure 3: X-ray diffraction pattern of (a) SLSG (b) RHA



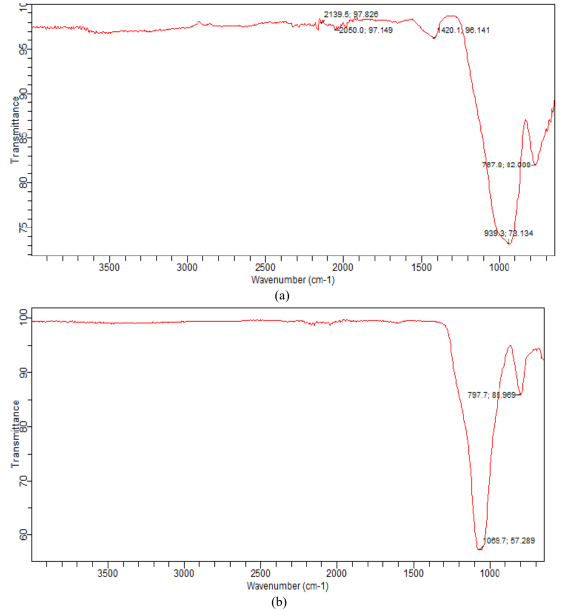


Figure 4: FTIR spectra of (a) SLSG (b) RHA

BET paran	neters	SLSG	RHA	
BET Total surface area $(m^2/g)$		1.9	22.3	
t-Plot micropore area $(m^2/g)$		1.3	0.2	
External surface	area $(m^2/g)$	0.60	22.2	
t-Plot micropore volume $(cm^3/g)$				
•		0.0007	0.0002	
Table 3. pH value		0.0007 ution synthesized from soda-l 170°C		
Table 3. pH value	es for sodium silicate sol	ution synthesized from soda-l	ime glass (SLSG)	
Table 3 pH value	es for sodium silicate sol 150°C	ution synthesized from soda-l 170°C	ime glass (SLSG) 200°C	

NaOH Conc.	150°C	170°C	200°C
3M	13.64	13.59	13.58
4M	13.82	13.84	13.77
5M	14.13	14.10	13.84

Silica source	NaOH Conc.	Synthesis Temp. (°C)	Viscosity at 25°C (Pa.s)
SLSG	3M	170	232
	4M	200	235
	5M	150	234
RHA	3M	150	314
	4M	200	314
	5M	170	305

Silica source	NaOH Conc.	Synthesis Temp (°C)	SiO2 (%)	Na2O (%)	H2O (%)	SiO <sub>2</sub> /Na <sub>2</sub> O	Product Name
SLSG	3M	170	30.83	16.26	52.9	1.89	Sodium disilicate
	4M	200	40.98	18.05	40.9	2.27	Sodium disilicate
	5M	150	30.16	14.38	55.5	2.09	Sodium disilicate
RHA	3M	150	28.64	12.32	59.0	2.32	Sodium disilicate
	4M	200	42.25	17.23	40.5	2.45	Sodium disilicate
	5M	170	36.97	18.44	44.6	2.00	Sodium disilicate

## 4. Conclusions

In this research, synthesis and characterization of sodium silicate solutions (water glass) from valorization of industrial wastes as silica source using hydrothermal method was investigated. Preliminary characterization of the selected wastes was conducted prior to their actual use as silica source in the synthesis of soluble sodium silicate solutions while physio-chemical analysis of the sodium silicate was also carried out.

The results showed that:

- Both rice husk ashes (RHA) and waste soda-lime glass (SLSG) consist of silica content which are high enough to justify their use as silica source in synthesis of sodium silicate as shown by the EDXRF
- The XRD results showed that both RHA and SLSG displayed amorphous silica broad peaks typical of amorphous silica band and which are known to be soluble as supported by literature; making them valuable source of silica for sodium silicate production.
- The FTIR results showed that both RHA and SLSG possessed typical Si O Si bonding and which justify their use as silica source in sodium silicate production.
- The Brunauer Emmett Teller (BET) analysis showed that RHA possessed high specific surface than SLSG which account for better reaction with NaOH resulting into more dissolution of silica in the waste unlike SLSG.
- Both temperature of synthesis and concentration (molarity) of NaOH contributed to the amount of SiO<sub>2</sub> dissolution into the obtained sodium silicate and Na<sub>2</sub>O present. However, both the SiO<sub>2</sub> and Na<sub>2</sub>O content and the SiO<sub>2</sub>/Na<sub>2</sub>O ratio satisfy most commercial grade sodium silicate solutions range of 1.6 3.3.
- The type of sodium silicate solutions obtained in this work is commercially known as sodium disilicate.
- Based on the results obtained it is therefore concluded that it is feasible to synthesize sodium silicate using waste soda-lime glass and rice husk as silica source instead of constituting environmental nuisance.

## **Conflict of Interest**

The authors declare no conflict of interest

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