Luliconazole Amalgamated Copper Nanoparticles for Enhanced Drug Delivery against Resistant Bacteria

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Submitted: 10-06-2023 Accepted: 21-06-2023

ABSTRACT: present investigation, luliconazole coupled copper nanoparticles were synthesized to overwhelm drug resistance in Staphylococcus aurous, responsible for dermal skin infections. Luliconazole, copper sulphate, Tri sodium citrate are used. Copper nanoparticles were produced by size reduction method by using Copper Sulphate and Tri sodium citrate. CuNPs were merged into gel base of carbopol which was formed by hot method. Carbapol gel shows no phase separation. The particle size of CuNPs was found to be 413.0± 30.2nm. The % EE was found to be 65.2%. On description unilamellar, sphereshaped vesicles with soft surface were detected under transmission electron microscopy. XRD of CuNPs was found out crystalline in nature. The zeta potential of nanoparticle shows less aggregation of particle with 15.1 ± 3.69 mV value. The amount of copper content was measured 5.9 microgram/10 mg of nanoparticles. In vitro release study of Cu NPs shows 96.5% release of drug and show effective antibacterial activity against Staphylococcus aureus. Conclusion: Luliconazole copper nanoparticles were produced and show excellent antibacterial activity upon S. aureus.

KEYWORDS: Copper Nanoparticle, Luliconazole, Carbopol gel, Antibacterial activity, Chemical reduction method, Copper sulphate.

I. INTRODUCTION

In the past decade, the cure of sickness has been consummate by administrating drugs to human body through different routes likes parental, oral, topical, sublingual, inhalation, rectal etc. The delivery of drug through topical denotes the application of drug onto the body employing vaginal, rectal, ophthalmic and skin as the route of administration. On human body, skin is widely used and accepted route for local application and constitutes the principal administration for local application. 1

The term topical drug delivery means administration for medicament containing formulation to the skin to openly care for the cutaneous manifestations of a common illness (e.g. psoriasis) or cutaneous disorders with the purpose of confining the pharmacological or other effect of the medicament inside or surface of skin.2 During recent years, a report will show interest in the synthesis and applications of various metallic nanoparticles due to their outstanding optical and electronic properties, especially copper, gold and silver nanoparticles. Copper nanoparticles (CuNPs) have gradually become an active area of research because of unique chemical, physical, electrical and optical properties, low cost, ease availability and exhibit good antibacterial properties. The prime advantage of CuNPs is their low cost and its availability compared to gold and nanoparticles, resulting in the sample synthesis and various applications of CuNPs.3, 4

Copper is easily available metal and one of the vital trace elements for mainly living creature. Copper was used as potential antimicrobial agent from ancient times. Copper and its complexes used as a disinfectants, antiviral as well as antibacterial from centuries. It is said that the enhanced antimicrobial activity of Cu-NPs due to their crystallographic surface structure and large surface to volume ratio compared with copper salts. Cu(OH)2 and CuSO4 are used as the conventional inorganic antibacterial agents. Also, complex copper species, aqueous copper solutions or copper containing polymers are used as antifungal compounds as well as antibacterial. At current, advancement in new antibacterial agents is essential due to steady raise of new bacterial strains resistant to the potent antibiotics. Substances with low molecular weight like copper nanoparticles generally inhibit the growth or kill a wide range of bacterium bacteria. Copper ions shows antimicrobial activity across a wide range of microorganisms, such as Salmonella enteric, Staphylococcus aureus, Campylobacter jejuni, Listeria monocytogenes and Escherichia coli. The surfaces of copper can be used to kill viruses, yeasts and bacteria hence copper known as "contact

International Journal of Pharmaceutical Research and Applications Volume 8, Issue 3 May-June 2023, pp. 2864-2875 www.ijprajournal.com ISSN: 2249-7781

killing".6,7 The Cu particles in nano range have been shown a antibacterial effect on the microbial cell functions in numerous ways, including electrostatic interaction between particles and gram negative bacteria cell wall, denaturation of the intracellular proteins and interaction with phosphorus- and sulfur containing compounds like DNA. The nanoparticles passed through bacteria cell membrane and then injurious for the vital enzymes of bacteria can be the primary mechanism of anti- microbial action in CuNPs.8,9

II. MATERIALS AND METHODS

The following chemicals were used: Luliconazole (Combiotic Pvt. Ltd. India), copper sulphate (Thermo Fisher Scientific, India), Tri sodium citrate (Nice chemicals, India.) Propylene glycol (S.D. Fine Chem. India), Methanol (Loba Chemie, India), Ethanol (Loba Chemie, India). Carbopol 940 (Lubrizol advanced material, Belgium), Methyl paraben (Lubrizol advanced material, Belgium), Propyl paraben (Lubrizol advanced material, Belgium), Triethanolamine (Nice chemicals, India).

Methods

Determination of melting point: In capillary tube, small amount of drug was added, and tube is sealed. The sealed tube was located in the melting point apparatus. The heat in the apparatus was slowly increased and the temperature at which whole drug gets melted was noted. DSC study of pure drug was conceded out on Shimadzu thermal analyzer DSC TA 60. The apparatus was calibrated using standard metal like high purity indium metal. The scans were conducted at heating rate of 10°C/min in nitrogen environment.

Solubility studies: The solubility study of Luliconazole was performed in methanol, ethanol, chloroform, acetone, distilled water, 0.1 N HCl, phosphate buffer solution pH 6.8, 7.4, individually by keeping the drug containing test tube on vortex mixture.

Preparation of standard curve in methanol: Accurately weighed 100 mg of luliconazole and transferred into 100 ml volumetric flask, make the volume up to 100 ml using methanol. From the above solution 10 ml was pipette out and transferred into 100 ml volumetric flask. The volume is made up with methanol in order to get standard stock solution containing 100 ppm. Form the above solution; a sequence of dilution (2, 4, 6, 8, 10 ppm) was diluted with the help of methanol. measured All dilutions were using

spectrophotometrically against blank of methanol at 220 nm for luliconazole. Absorbance of drug at different concentrations was calculated and graph was plotted.10

Infrared spectroscopic analysis: The FTIR spectrums of luliconazole, carbopol and mixture of luliconazole, carbopol were recorded on IR spectrophotometer. All the samples are free from moisture. Infrared spectrum was recorded in the 4000-400 cm⁻¹ regions (Bruker).

COPPER NANOPARTICLES SYNTHESIS: Cu nanoparticles were synthesized using size reducing agent like tri sodium citrate. Copper sulphate and trisodium citrate employed as initial substances in the development of copper NPs. All solutions were prepared in distilled water. Make 0.001 M CuSO4 solution with distilled water, take 40 ml from this solution in beaker and heated the solution to boil. In above solution, 10mL of 1% trisodium citrate was mixed drop wise. The mixture was heated under continuous magnetic stirring for 30 minutes. The mix solution was them cooled near room temperature. The reaction was allowed to take place for 24 hr. Accurately weighed 2g luliconazole was dissolved in methanol and added to copper nanoparticles.11

PREPARATION OF THE CARBOPOL GEL:

At low concentration, carbopol 940 forms very good flexible transparent gel. The gel base of 2% was prepared by scattering 2 g carbopol 940 in 86 ml warm distilled water. Accurately weighed 0.6 g propyl paraben and dissolved in ethanol. Accurately weighed 0.3 g methyl paraben and dissolved in 15 ml of propylene glycol. Stirred the mixture unless gelling occurred and then mixture was neutralized with the help of 50% (w/w) triethanolamine. Triethanolamine was added drop by drop to maintain the pH between 6-7.12

The nanoparticle formulation containing drug was slowly added in carbopol 940 gel base and mixed with the help of stirrer for 5 min continue stirring.

EVALUATION OF NANOPARTICLES

Drug entrapment efficiency: Take 5 ml formulation and diluted the formulation up to 8 ml with distilled water and centrifuged the diluted formulation at 15,000 rpm at 4°C for 45 min using a cooling centrifuge. The sediment and supernatant were restored after centrifugation, their volume was calculated. Then sediment was break down through n-

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International Journal of Pharmaceutical Research and Applications

Volume 8, Issue 3 May-June 2023, pp: 2864-2875 www.ijprajournal.com ISSN: 2249-7781

propanol and filtered using a $0.45~\mu m$ nylon filter. The concentration of luliconazole in the sediment and supernatant was examined by UV- spectrophotometer at 220 nm. The % entrapment efficiency was estimated.

Percentage Entrapment efficiency Amo unt of entrapped drug recovered

Total amount of drug

- Nanoparticle shape: Transmission electron microscopy (Philips Technai electron microscope, Netherlands) was used for the forecast of nanoparticle. At room temperature, sample was dried and vesicular were forecast under microscopy working at an acceleration voltage of 200 KV for 5 min.
- Particle size estimation: Dynamic light scattering method was used for the determination of copper nanoparticles, using a computerized inspection system (Malvern Zetasizer Nano-ZS, Malvern, U.K.). For the measurement of size, copper nanoparticle solution was attenuated with distilled water and implement in cuvettes of zetasizer.13
- **Zeta potential measurement:** Physical property like zeta potential which describe the net surface charge of copper nanoparticles. The stability criteria of CuNPs are measured when the zeta potential values ranges from higher than +30 mV to lower than 30 mV.¹⁴
- X-ray diffraction: 1 ml of the copper nanoparticle solution was extend on a glass slide and dried at 40°C in an oven. The Phillips Xpert a glass slide and dried at 40°C in an oven. The Phillips Xpert Pro Diffractometer were recorded the spectra running at 40 kV and 30 mA.15
- Copper content determination:
 Determination the copper (II) ions, take 200 ml
 of tap water in the beaker. Water is evaporated
 up to 50 ml Solution is transferred into a
 volumetric flask and the determination is
 performed. To different volumes of water, the
 solution containing copper are added and the
 solution is was brought up to the mark by
 mixture of acetate buffer. The absorbance is
 measured at a wavelength of 520 nm.16

Physical evaluation of nanoparticle gel pH measurement of the nanoparticle gel: 1 gm luliconazole coupled nanoparticle gel base was mixed in 100 ml beaker containing distilled water. After that pH electrode was deep in beaker and readings were reported from digital pH meter.

- Viscosity study: The viscosity of copper nanoparticle was measured in Brookfield instrumentation by selecting appropriate spindle and rpm. In 50 ml beaker, 50 g of formulation was added which was set till spindle channel was drenched and set rpm. Reading pointed out over three minutes.
- **Spreadability:** Spreadability term denote a area is required to which gel willingly fall on appliance to skin or affected part.17,18 It was calculated through formulation:

 $S=M X \frac{L}{T}$

Where T: time taken to separate the slides L: length of slides

M: wt. tied to upper slide

- Extrudability study: The extrudability of luliconazole coupled nanoparticle gel was considered by stuffing nanoparticle gel in the foldable tubes. Determination in words of weight in grams, 10 sec required to extrude a 0.5 cm ribbon of gel.
- **Percentage yield:** Percentage yield was calculated by the formula.

 $Percentage\ yield = \ \frac{Practicle\ yield}{Theortical\ yield}\ X\ 100$

- **Grittiness and homogeneity:** A tiny amount of nanoparticle gel was squeezed in the middle of index finger and the thumb. Uniformity of the nanoparticle gel is observed, any crude particles visible on fingers. 19,20
- In vitro release studies: Vertical Franz diffusion cell apparatus was employed for the in vitro absorption studies. It contain donor as well as receptor chamber that is filled with PBS. The donor chamber is filled and the permeation of solute through the membrane is monitored at different interval of time. Episodic sampling from the receptor chamber was collected and measured. The jacketed cell personified is stirred during experiment at 500 rpm using a magnetic agitator.21

Drug release kinetics: The kinetic of drug release was calculated by various kinetic models as zero order release kinetics plot, first order release kinetics plot, korsmeyer-peppas release kinetics plot and higuchi release kinetics plot. To study profile release kinetics of the copper nanoparticle figures obtained from in-vitro release profile were plotted for various kinetic models. The finest fit model was set by the value of R2 close to 1.22



Volume 8, Issue 3 May-June 2023, pp: 2864-2875 www.ijprajournal.com ISSN: 2249-7781

Antimicrobial activity studies: Antimicrobial activity has been assayed against bacteria by using agar diffusion method. The antibiotics action of drug is as oleic of its capability to growth inhibition of bacto nutrient agar or broth. Cup-plate method shall be used for the consideration of bacterial inhibition. In experiment, discs of average diameter were prepared in the bacto agar nutrient medium, containing standard bacterial inoculums. The test samples are injected in the disc and the diameter of the zone of inhibition was evaluated. All the test samples were evaluated for antibiotics activity against Staphylococcus aureus (gram positive).23

III. RESULT AND DISCUSSION

- **Melting point determination:** Melting point of pure luliconazole outcome at 78°C. Melting Point outcome three times and mean was noted.
- **Solubility studies:** Luliconazole was found to be soluble in acetone, methanol, 0.1 N HCL, distilled water, ethanol, chloroform, PBS of 6.8, 7.4.

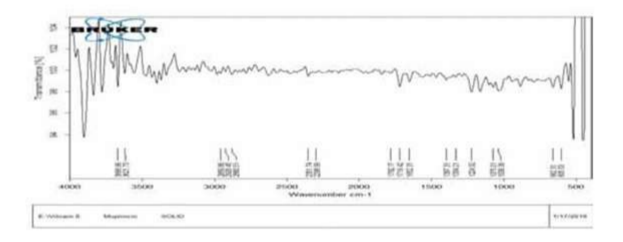


Figure 1: FTIR of Luliconazole

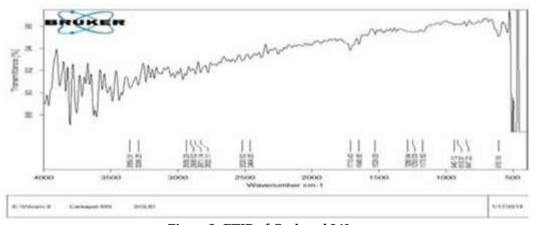


Figure 2: FTIR of Carbopol 940

FTIR analysis: FT-IR analysis discovered that there was no interaction between the luliconazole and physical mixture as per given in pharmacopoeia. In the present investigation, it has been observed that there are no chemical and physical interactions because of some bond

formation between luliconazole and physical mixture. Hence luliconazole drug was authentic and free from impurities. FTIR of Luliconazole, Carbapol and physical mixture of Luliconazole and Cabool respectively).



Figure 3: FTIR of Physical mixture.

Calibration curve of luliconazole: The graph obeyed beer lamberts law in this selected concentration range.

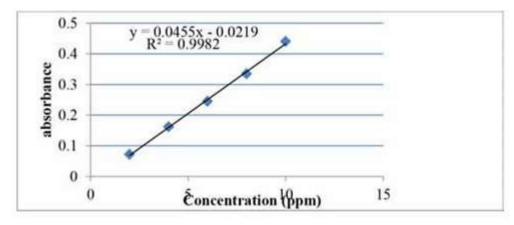


Figure 4: Standard calibration curve for different dilution of luliconazole at 220 nm.

Straight line was observed to be $y=0.045x\ \Box\ 0.021$ with correlation coefficient 0.998 shown in Figure 5.

EVALUATION OF NANOPARTICLES

• **Drug entrapment efficiency:** % E.E. of drug was found to be 65.2%.

• Transmission electron microscopy (TEM): Formulation was subjected for TEM to obtain, image of nanoparticles on scale bar of 200 nm with magnification 13.0x4000. On description unilamellar, spherical vesicles with smooth surface were noticed under transmission electron microscopy (TEM) shown in Figure 5.

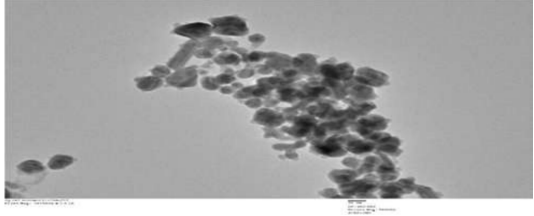


Figure 5: TEM of copper nanoparticles



Zeta potential: The zeta potential of copper nanoparticle shows in Figure 7, less aggregation of particle with 15.1 ± 3.69 mV values.

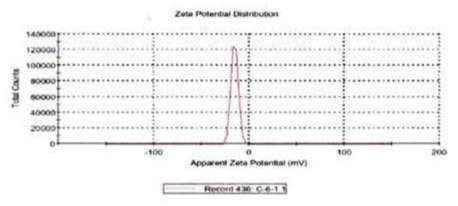


Figure 6: Zeta potential Copper nanoparticles

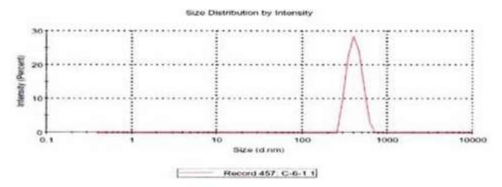


Figure 7: Particle size Copper nanoparticle

- **Particle size measurement:** Dynamic Light Scattering of copper nanoparticles; the average size obtained was of 413.0 ± 30.2 nm with a narrow size distribution.
- X-ray diffraction (XRD): The crystal

structure and phase composition of synthesized copper nanoparticles is analyzed by XRD. The diffraction data exhibits that the copper nanoparticles have crystalline structure.

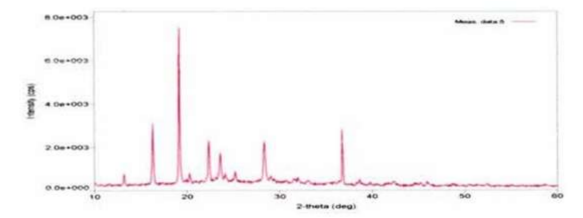


Figure 8: XRD spectra of copper nanoparticles



Volume 8, Issue 3 May-June 2023, pp: 2864-2875 www.ijprajournal.com ISSN: 2249-7781

- Copper content determination: The amount of copper content was measured 5.9 microgram/10 mg of nanoparticles.
- Physical Evaluations of Nanoparticle Gel Organoleptic characteristics: Color is pale yellow, Odor is characteristic, Phase separation is no.
- **Determination of pH of nanoparticle gel**: The pH of nanoparticle gel was recognized as 7.1
- **Viscosity:** The viscosity of carbopol 940 gel base and nanoparticle gel by Brookfield instrumentation was recognized 73,200 and 72,300 cP respectively.
- **Spreadability:** The spreadability of luliconazole gel coupled copper nanoparticle was recognized 13.29 g.cm²/sec. The results demonstrated that gel was effective.
- Extrudability analyzed: The extrudability of luliconazole gel coupled copper nanoparticle was recognized positive. Positive extrudability showed the better application of gel.

- **Percentage yield**: The % yield of luliconazole gel coupled copper nanoparticle was carried out 95.78%.
- Homogeneity and grittiness: Luliconazole gel coupled copper nanoparticle was recognized homogeneous and no grittiness was indicated.
- In vitro release profile: Franz diffusion cell apparatus was used for the in vitro release profile. The drug release profile of the luliconazole coupled copper nanoparticles is presented in Figure 10. In vitro release profile was performed24-41 to determine amount of drug release at different interval of time. The cumulative drug releases from nanoparticle reach 96.5% in 48 hr.

Drug release kinetics: The kinetics for drug release of luliconazole coupled Cu NPs was carried out for different models.

Zero Order release kinetics plot: Plot the graph % cdr Vs time.

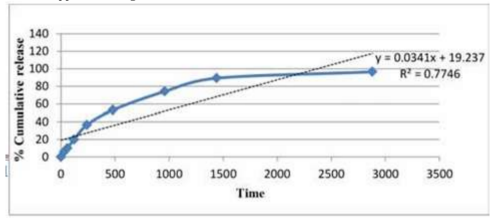


Figure 9: Zero Order release kinetics plot.

• **First Order release kinetics Plot:** Graph was prepared between log % cumulative drugs remaining Vs time.

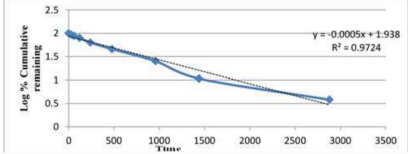


Figure 10: First Order release kinetics Plot.

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• **Higuchi's Model release kinetics:** Graph was prepared between % cdr Vs square root of time (Graph shown in Figure 12).

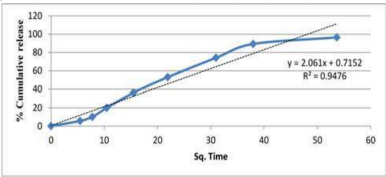


Figure 11: Higuchi plot for release kinetics plot.

• Korsmeyer-Peppas Model release kinetics: Graph was prepared between log % cdr Vs log time (Graph shown in Figure 14). Some kinetic models describing drug release from modified released dosage forms. The model release data by correlation coefficient. The correlation

coefficient value was used as criteria to choose the best model to explain the drug release. From these values, it was observed that the peppas model will be fitted best model with R^2 value of 0.980.

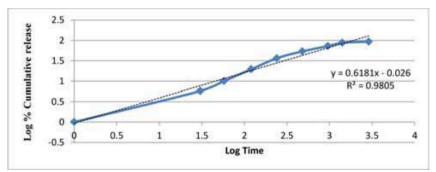


Figure 12: Peppas model release kinetics plot.

Table 1: Kinetics of drug release

Plot	K ₀	R ²
Zero order	0.077	0.774
First order	0.000	0.972
Higuchi	4.75	0.947
Peppas	1.41	0.980

ANTIMICROBIAL ACTIVITY: Antimicrobial activity was determined by cup plate method on S. aureus. Antimicrobial activity of pure drug shown

in Figure 15 and Antimicrobial activity of copper nanoparticles gel containing luliconazole.



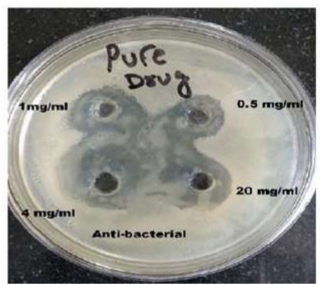


Figure 13: Antibacterial activity of pure drug luliconazole

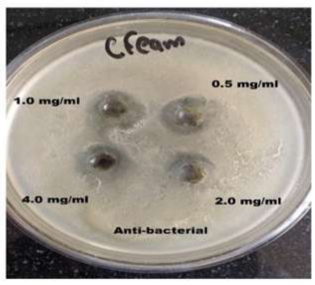


Figure 14: Antibacterial activity of copper Nanoparticlegel containing luliconazole.

Table 2: Compare Antimicrobial activity by cup-plate method.

Table 2. Compare Anumerobial activity by cup-plate method:			
Concentration (µg/ml)	Pure Luliconazole	Formulation	
0.5	9 ± 1 mm	6± 2 mm	
1	12 ± 2 mm	9± 2 mm	
4	17 ± 1 mm	$13 \pm 3 \text{ mm}$	
20	18± 3 mm	17 ± 1 mm	
30	18± 2 mm	18 ± 2mm	
40	19± 2 mm	18 ± 1 mm	

IV. CONCLUSION:

In present investigation, luliconazole coupled copper nanoparticles were synthesized to overwhelm drug resistance in Staphylococcus

aureus, responsible for dermal skin infections. So, prepared and evaluate the luliconazole coupled Cu NPs to get the formulation with increased antibacterial activity and suit for topical



Volume 8, Issue 3 May-June 2023, pp: 2864-2875 www.ijprajournal.com ISSN: 2249-7781

application. The Cu NPs containing luliconazole were prepared by chemical reduction method and evaluated. Based on R2 value the formulation followed the Pappas model. Luliconazole containing Cu NPs based gel displayed superior efficacy against S. aureus owning to prolonged release as compared to pure drug.

CONFLICT OF INTEREST: The authors declare no conflict of interest.

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Volume 8, Issue 3 May-June 2023, pp: 2864-2875 www.ijprajournal.com ISSN: 2249-7781

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