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Aim: Diabetic (type-2) is a metabolic disease characterized by increased blood glucose level from the normal level. In the present study, apigenin (AG) loaded lipid vesicles (bilosomes: BIL) was prepared, optimized and evaluated for the oral therapeutic efficacy.

Experimental: AG-BIL was prepared by a thin-film evaporation method using cholesterol, span 60 and sodium deoxycholate. The prepared formulation was optimized by 3-factor and 3-level Box-Behnken design using particle size, entrapment efficiency and drug release as a response. The selected formulation further evaluated for ex-vivo permeation, in vivo pharmacokinetic and pharmacodynamics study.

Results: The optimized AG bilosomes (AG-BILopt) has shown the vesicle size 183.25 ± 2.43 nm, entrapment efficiency 81.67 ± 4.87%. TEM image showed a spherical shape vesicle with sharp boundaries. The drug release study revealed a significant enhancement in AG release (79.45 ± 4.18%) from AG-BILopt as compared to free AG-dispersion (25.47 ± 3.64%). The permeation and pharmacokinetic studies result revealed 4.49 times higher flux and 4.67 folds higher AUC0-t than free AG-dispersion. The antidiabetic activity results showed significant (P < 0.05) enhancement in therapeutic efficacy than free AG-dispersion. The results also showed marked improvement in biochemical parameters.

Conclusion: Our findings suggested, the prepared apigenin loaded bilosomes was found to be an efficient delivery in the therapeutic efficacy in diabetes.

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flavones category. It is found in various natural sources like vegetables and fruits (Funakoshi-Tago et al., 2011). There are number of pharmacological activities like anti-diabetic (Choi et al., 2014), anticancer (Lu et al., 2020), anti-oxidant (Darabi et al., 2020) and anti-inflammatory activity (Wang et al., 2020) have been reported. It has poor aqueous solubility and belongs to BCS-II class category (Huang et al., 2016). Due to poor aqueous solubility, their bioavailability is limited (~7.1%) and led to poor therapeutic efficacy after single-dose oral administration in the rat (Perez-Moral et al., 2018, Elhennawy and Lin, 2018).

There are several nano-sized lipid vesicles of poorly soluble drugs have been reported to enhance the solubility and bioavailability (Mishra et al., 2016; Paudel et al., 2017; Freag et al., 2018; Pauli et al., 2019; Sze et al., 2019; Rizwanullah et al., 2017). Bilosomes (BIL) is an elastic nano-size vesicle consists of lipid, surfactant and bile salt (Parashar et al., 2019). It is structurally similar to liposome (Yang et al., 2019a, 2019b), and protect the drug molecule from the enzymatic degradation into GIT (Wilkhu et al., 2013). The bile salt present in intestine (GIT) limits the capabilities of the conventional vesicle by causing lysis of vesicle membrane and led to early release of entrapped drug or molecule before reaching to the site of action (Naguib et al., 2020). To overcome the limitations of conventional nano-vesicular drug delivery systems, the application of bilosomes (bile salt stabilized nano-vesicle) as a delivery system has been widely accepted. It is a nanosized vesicle prepared with the addition of bile salt into a lipid bilayer. The bile salt is biologically compatible and having no toxicity (Nuruunabi et al., 2016; Ahmad et al., 2017). It has assimilation enhancing ability and acts as solubilizing and permeation enhancing agent. It may enhance the drug permeability and bioavailability of poorly soluble therapeutics (Deng and Bae, 2020). Bile salts absorbed through apical sodium-dependent bile acid transporter (ASBT) in the GIT, and may also act as an enhancer in the oral delivery (Nurunnabi et al., 2016). The different type of bile salts such as sodium deoxycholate, sodium glycolate and sodium taurocholate were used to formulate BIL. Among them, sodium deoxycholate is most commonly used due to its nontoxic and high permeation enhancing capacity (Niuruunabi et al., 2016; Ahmad et al., 2017). It has assimilation enhancing ability and acts as solubilizing and permeation enhancing agent. AG loaded bilosomes were prepared by the thin-film hydration method with slight modification (Saifi et al., 2020). The weighed quantity of cholesterol (CHO), surfactant (Span 60), and Apigenin (AG) were taken in a round bottom flask and dissolved in organic solvent (10 ml). The round bottom flask attached to the rotary evaporator (IKA, RV-3 V, Germany) and the organic solvent was evaporated at 55 °C under reduced pressure. After complete removal of the solvent, the flask was kept overnight in desiccator to remove the residual solvent. The dry thin film was hydrated with bile salt (Sodium deoxycholate, SDC) containing distilled water for 45 min. Finally, AG loaded bilosomes (BIL) was collected and stored for further characterization.

4. Formulation optimization

Box-Behnken design (BBD) was used to optimize the prepared AG BIL. This design was used most commonly because it gives lesser number of runs with five centre point (Ameeduzzafar et al., 2014; Khan et al., 2018). The independent variables Cholesterol (A, CHO), surfactant (B, span 60) and bile salt (C, sodium deoxycholate) were used at three level (low, medium, high) as independent variables. There effects were assessed on vesicle size (Y1), entrapment efficiency (Y2, EE) and drug release (Y3) as shown in Table 1. The design showed seventeen experimental runs with five centre point (same composition) and the detail compositions were expressed in Table 2.

4.1. Characterization

4.1.1. Vesicle characterization

The vesicle size, PDI, and zeta potential of the prepared AG-BIL were determined by size analyzer (Malvern zeta sizer, Malvern., USA). The sample of AG-BIL (0.1 ml) were diluted 100-fold and transferred into cuvette. The sample were scanned at 90 ° scatter-angle at room temperature. The morphology of the selected AG-BIL was performed by using high resolution TEM. The sample was stained and evaluated under the electron microscope to evaluate the morphology.

4.1.2. Entrapment efficiency (EE)

The entrapped AG into BIL was estimated by ultracentrifugation method (Ameeduzzafar et al., 2014). The appropriate volume of BIL was taken into centrifugation tube and centrifuged at 18000 rpm for 30 min at 4 °C (Sigma 3-16KL, Osterodeam Harz, Germany). The supernatant was separated and AG concentration was analyzed by using UV–Vis-spectrophotometer (Genesys10S UV–Vis, Thermo Scientific, USA) at 269 nm with appropriate dilution.

3. Methods

3.1. Formulation of AG bilosomes

AG loaded bilosomes were prepared by the thin-film hydration method with slight modification (Saifi et al., 2020). The weighed quantity of cholesterol (CHO), surfactant (Span 60), and Apigenin (AG) were taken in a round bottom flask and dissolved in organic solvent (10 ml). The round bottom flask attached to the rotary evaporator (IKA, RV-3 V, Germany) and the organic solvent was evaporated at 55 °C under reduced pressure. After complete removal of the solvent, the flask was kept overnight in desiccator to remove the residual solvent. The dry thin film was hydrated with bile salt (Sodium deoxycholate, SDC) containing distilled water for 45 min. Finally, AG loaded bilosomes (BIL) was collected and stored for further characterization.

2. Materials and methods

2.1. Materials

Apigenin was procured from “Beijing Mesochem Technology Co. Pvt. Ltd. (Beijing, China)”. Span 60, diethyl ether, chloroform, HPLC grade ethyl acetate, methanol, acetonitrile and water were procured from Sigma Aldrich (St Louis, MO63103, USA). Formic acid was procured from SD-fine chemical (Mumbai, India). Cholesterol and Sodium deoxycholate were obtained from Thermo Fisher Scientific, India. Dialysis bag (MW-cut off 12000) was purchased from the Sigma Aldrich (St Louis, MO63103, USA).

| Table 1 | Independent variables and responses used for optimization of bilosomes. |
| --- | --- |
| Factor | Level |
| Independent variables | Low (−1) | Medium (0) | High (+1) |
| A: Cholesterol (CHO, %) | 10 | 20 | 30 |
| B: Surfactant (Span 60, %) | 50 | 60 | 70 |
| C: Bile Salt (Sodium deoxycholate, SDC) | 10 | 15 | 20 |
| Responses | Y1: Vesicle size (nm) | Optimum | Y2: Entrapment efficiency (%) | Maximize |
| | Y3: Drug release (%) | Maximize |
The study was performed in triplicate and % EE was calculated by below formula:

\[
\text{EE} (%) = \frac{\text{Total drug} - \text{Free drug in supernatant}}{\text{Total drug}} \times 100
\]

4.1.3. Drug release

The release study of AG-BIL formulations were studied by dialysis bag method (Ameeduzzafar et al., 2020). The dialysis bag (mw ~ 12000, St Louis, MO63103, USA) was prepared for the study as per the standard procedure and then soaked with release media (simulated intestinal fluid, SIF). The appropriate volume (~5 mg of AG) was filled into dialysis bag and tied from both side and immersed into released media (900 ml) with tween 80 (0.3% v/v). The release media was rotated at 100 rpm and temperature was maintained at 37 ± 0.5 °C during the study. At predetermine time intervals (30 min, 1, 2, 4, 6, 8, 12 h) a fixed volume (5 ml) sample was withdrawn and replaced with fresh release media. In similar intervals (30 min, 1, 2, 4, 6, 8, 12 h) a fixed volume (5 ml) sample was withdrawn and replaced with fresh release media. The collected samples were filtered, diluted and drug concentration was determined using UV spectrophotometer at 269 nm. The study was performed in triplicate and % EE was calculated by reported HPLC method (Cai et al., 2006). The permeation flux and apparent permeability was calculated for both the sample to compare the enhancement.

5. Biological study

5.1. Animal handling

The comparative preclinical evaluation of AG-BILopt and free AG dispersion was done on Wistar male albino rat (200–250 gm). The study protocol was approved (04/02/41) by institutional animal ethical committee of Jouf University, Aljouf, Saudi Arabia. The animal was procured from animal house and kept in standard environmental condition (12 h dark/light cycle). The animals were fed with standard rich fat diet contain normal pellet diet, cholesterol, casein protein, vitamin, coconut oil, sucrose, fructose, sodium chloride and dl- methionine for 15 days. The blood glucose level (BGL) of each animal was measured using glucometer (Accu-Check, Roche, Germany) at zero time and further after treatment with AG-BILopt and free AG dispersion of same dose. The animals were divided into four groups (group 1 considered as normal control (NC); group 2 taken as diabetic control (DC), group 3 and group 4 received Free AG dispersion and AG-BILopt. Each treatment group has six rats as shown in Table 4.

5.2. Induction of diabetes

Streptozotocin (STZ) with high fat diet model was used for induction of hyperglycaemia (Type 2 diabetes) in rats. STZ (0.1 M) was freshly prepared in citrate buffer (pH 4.5) and administered single low dose (35 mg/kg) intraperitoneally (Guo et al., 2018; Ameeduzzafar et al., 2019). The animals were kept for 72 h to stabilize the BGL and after that the fasting BGL was measured. The animal having ≥ 200 mg/dl fasting BGL were marked as hyperglycaemic animals and considered for further experiments.

5.3. Pharmacokinetic study

The pharmacokinetic study of the prepared AG-BILopt and free AG dispersion was performed in Wistar albino rats. The AG-BILopt and free AG-dispersion (~60 mg/kg of AG) was administered into group 3 and group 4 animals (Ding et al., 2014). At predetermine time interval (0, 0.5, 1, 2, 6, 12, and 24 h) animals were anesthetized using diethyl ether inhaler and the blood was
collected into EDTA tube. The blood was centrifuged at 5000 rpm for 15 min and plasma was separated. AG was extracted from the plasma by liquid phase extraction technique. Formic acid (2% v/v) was mixed with plasma and vortexed for 10 min. Then ethyl acetate (2 ml) was added, mixed and centrifuged for 15 min (3000 rpm) at 4°C. The supernatant was collected and dried it under nitrogen evaporator. The sample reconstituted with methanol (500 ml), filtered by membrane filter (0.25 mm) and injected (20 ml) into HPLC system (Auto-sampler) for estimation of AG concentration. The different pharmacokinetic parameters i.e., Cmax, Tmax, T 1/2, AUC0-t, AUC0-1, Kel and AUMC0–24 was determined by using of software (Excel add-on PK solver).

5.4. Evaluation of hyperglycaemic activity

The prepared AG-Bilopt and free AG-dispersion formulations were orally administered one time a day in appropriate dose (60 mg/kg) in the treatment animal group 3 and group 4 using oral feeding needle (Ding et al., 2014). At predetermined time interval one drop of blood was taken directly into glucometer strip from rat tail. After some time (5 s), BGL displayed on the screen of glucometer and noted. The reduction (%) BGL was calculated by given formula for both groups

\[
\text{BGL reduction (\%)} = \frac{BGL_t - t_0 - BGL_{t-1}}{BGL_{t-1} - t_0} \times 100
\]

5.5. Biochemical evaluation

The biochemical parameters were evaluated at the end of study. The collected blood sample was kept aside for some time and plasma was separated by centrifuging it at 3000 rpm for 15 min. The blood plasma was stored in refrigerator for the biochemical evaluation. The parameters like lipid profile, urea, uric acid, serum total protein, creatinine, Serum glutamic–pyruvic transaminase, and serum glutamic oxaloacetic transaminase were evaluated using commercially available standard kits by PAP method.

5.6. Statistical analysis

The study was performed in triplicate and data was represented as mean with standard deviation. The graph pad prism was employed to determine the statistical calculation. The P value < 0.05 considered as statistically significant.

6. Result and discussion

6.1. Optimization

The design showed total seventeen experimental runs from BBD with five centre point (same composition) as represented in Table 2. These experimental results were fitted into the software for optimization. The vesicle size, entrainment efficiency and drug release of AG-Bil formulations were found in the range of 118.93–267.32 nm, 71.62–93.04 %, and 76.34–92.87 %, respectively. The effect of independent variable on dependent variable were assessed graphically i.e., 3D response surface plot (Figs. 1A-C). It helps to express the effect of more than one variable over one response at one time. The results of each dependent variable were fitted into various experimental models to evaluate the best fit model. The best model was found to be quadratic model and the maximum regression value (R²) was found than other models. The actual
The experimental and predicted (software) value of each response was found to be very close to each other (Table 3). The effect was also expressed by mathematical polynomial equation and by numerical value of the actual vs predicted value graph (Fig. 2). The analysis of variance for each response was calculated and the value was found to be $P < 0.0001$ (Table 4).

**Fig. 1A.** 3D response surface plot showing effect of independent variable over the vesicle size.

**Fig. 1B.** 3D response surface plot showing effect of independent variable over the entrapment efficiency.
6.2. Effect of cholesterol, span 60 and sodium deoxycholate on vesicle size

The vesicle size of all the prepared AG-BIL formulations were found in range of 118.93–267.32 nm (Table 2). The polynomial equation and 3D response surface graph (Fig. 1A) showed the effect of independent variables on the size. Cholesterol (CHO) exhibited positive effect on vesicle size. As the CHO concentration increases the vesicle size increases. CHO hindered the compact packing of lipid vesicle and subsequently higher aqueous phases inside the vesicle (Fig. 1C).

**Fig. 1C.** 3D response surface plot showing effect of independent variable over the drug release.

**Fig. 2.** Actual and predicted value graph of A) Vesicle size, B) Entrapment efficiency, C) Drug release (%).
bilosome vesicle and lead to increase in vesicle size (Ameeduzzafar et al., 2020). The surfactant showed the negative effect on vesicle size. The increase in surfactant concentration led to reduction in interfacial tension between CHO and aqueous phase and size decreases. The third variable bile salt also exhibited the negative effect on vesicle size. As the bile salt concentration increases the vesicle size decreases due to reduced surface tension as well as flexibility of BIL. But at high concentration it tends to form aggregates itself (Yang et al., 2019a, 2019b). The computer-generated polynomial equation of quadratic model for vesicle size is given below (Eq. (1))

\[
Vesicle \ Size \ (Y_1) = + 183.67 + 47.36A - 26.57B \\
- 9.57C - 8.23AB + 1.37AC \\
+ 9.12BC - 1.85 A^2 + 3.07 B^2 \\
+ 9.85 C^2.
\] (1)

where the A, B, C, AB, AC, BC, A^2, B^2, and C^2 are the significant model term (P < 0.05). The model F value was found to be 5811.20 suggested that model is significant (P < 0.0001). The F-value Lack of fit is 0.15, suggested that not significant expressed that model was well fitted. The predicted R^2 of 0.9996 is in reasonable agreement with adjusted R^2 of 0.9997 and the adequate precision was found to be > 4 (276.62), represented the model have adequate signal (Table 3). The actual and predicted value was expressed by graphical presentation (Fig. 2A).

6.3. Effect of cholesterol, span 60 and sodium deoxycholate on entrapment efficiency

The entrapment efficiency was found in range of 71.62–93.04 % (Table 2). The effect of independent variables on response was represented by 3D response surface graph (Fig. 1B). From the graph and below polynomial equation, it was observed that CHO (A)
Sodium deoxycholate concentration increases the entrapment efficiency because SDC have surface active property and incorporated into bilayer membrane surface, increases the flexibility of lipid membrane and increasing the solubility of drug in the lipid membrane hence increased the entrapment efficiency. The computer generated polynomial equation written below to interpret the results.

\[
EE (Y_2) = +81.67 + 8.16A + 3.57B + 0.33C \\
- 0.95 AB - 0.68AC + 1.23BC - 0.14 A^2 \\
+ 0.71B^2 - 0.67C^2 
\]

(2)

In this polynomial equation, the model term A, B, C, AB, AC, BC, B^2, and C^2 are significant (P < 0.05, i.e., significant effect on the entrapment efficiency) and A^2 has shown non-significant model term (P > 0.05). The positive and negative sign of equation indicates the synergistic and antagonistic effect to the tested responses. The model F-value (868.6) implies that model is significant (P < 0.0001) and well fitted. The lack of fit value is very low (F = 0.9) and suggested that lack of fit is non-significant and as well as fitted for quadratic model. The predicted R^2 of 0.9996 is in reasonable agreement with the adjusted R^2. The adequate precision is 276.619 (<4), and represented the model has adequate signal. The actual and predicted value was expressed by graphical presentation (Fig. 2B).

6.4. Effect of cholesterol, span 60 and sodium deoxycholate on drug release

The drug release of all experimental runs (actual value) is 76.34–92.87% as shown in Table 2. The 3D and contour plot were generated and explained the effect of two factors on one response (Fig. 1C). CHO (A) exhibited negative effect on drug release, as the concentration of CHO (A) increases the drug release gradually decreases. The increases in CHO concentration, the vesicle wall become stiff and impede drug release from the vesicle. The second factor, surfactant (B) has shown the negative effect on drug release. At higher surfactant concentration, AG release from the BIL decreased due to the increased viscosity. The other reason for this type of release behaviour of AG into the dissolution media due to higher transition temperature and low HLB value of surfactant (span 60, B). The third factor bile salt (SDC, C) exhibited positive effect on drug release. The increases in bile salt concentration lead to increased drug release. On increasing the bile salt (SDC) concentration, it incorporated into core of BIL membrane and increases the flexibility. It gives increased release of AG from the formulation but at higher concentration it the drug release decrease.

The software generated polynomial equation of drug release was expressed by Eq. (3)

\[
\text{Drug Release (Y_3)} = +86.47 - 3.46A - 4.81B + 3.14C \\
+ 0.46 AB + 5.000E - 0.03 AC \\
+ 0.27BC - 1.41A^2 - 0.8 B^2 \\
- 0.34C^2
\]

(3)

The polynomial equation showed the A, B, C AB, A^2, and B^2 are significant (P < 0.05) affected on drug releases and remaining AC, BC, and C^2 are insignificantly (P > 0.05) effects on drug release. The positive and negative sign represent the synergistic and antagonistic effect on the drug release. The model F-Value of quadratic model for drug release was found to 393.04 suggested that model is significant (P < 0.05) and well fitted. The lack of fit is non-significant (P > 0.05) and represents ideal for the model. The predicted R^2 (0.9943) is in reasonable agreement with adjusted
The predicted AG bilosomes (AG-BILopt) has shown the vesicle size of 183.25 ± 2.43 nm and encapsulation efficiency of 81.67 ± 3.87% using the composition CHO (15.5%), surfactant (span 60, 70.2%), bile salt (SDC, 12.4%) and used for further study.

6.5. Vesicle characterization

The vesicle size of prepared bilosomes were found in the range of 118.93–267.32 nm (Table 2). The optimized formulation (AG-BILopt) showed vesicle size of 183.25 ± 2.43 nm and graphically represented in Fig. 3A. PDI of AG-BILopt was found to be 0.42 (<0.5), and it indicates homogeneous distribution of bilosomes. The zeta potential value (-31.8 mV) of AG-BILopt indicates higher stability. TEM was used to evaluate the morphology of AG-BILopt and the sample exhibited spherical shape (Fig. 3B).

6.6. Entrainment efficiency (%)

The entrainment efficiency of prepared AG-BILs were determined by centrifugation method and result showed in the range of 71.62–93.04% (Table 2). The optimized formulation AG-BILopt has shown 81.67 ± 3.87% encapsulation efficiency.

6.7. Differential scanning calorimetry

DSC Thermogram of AG, CHO, sodium deoxycholate and AG-BILopt were analysed and depicted in Fig. 4A-D. The thermogram of AG, CHO, and sodium deoxycholate exhibited its characteristic peak at 356 °C, 150 °C and 260 °C, respectively (Fig. 4 A-C). The thermogram of AG-BILopt showed only one characteristic endothermic peak at 145 °C (Fig. 4D), which is closer to the melting point of CHO. There was no any other characteristic peak AG was found in thermogram, indicates that AG was completely encapsulated into lipid bilayer matrix.

6.8. In-vitro release study

The comparative release study of AG from the prepared AG-BILopt and free AG dispersion was expressed graphically in Fig. 5. AG-BILopt showed biphasic release behaviour, an initial burst release (39.58 ± 2.32% in 2 h) and later prolonged release (79.45 ± 3.18%) was found for 12 h. The burst release was found due to the release of adsorb AG from the BIL surface whereas, prolonged release is due to more affinity of AG toward the hydrophobic part of bilosomes (Jain et al., 2014). The free AG dispersion exhibited poor drug release of 25.47 ± 1.64% than AG-BILs formulation. The release data was fitted into different release kinetic model for determination of best fit model on the basis of maximum regression co-efficient value ($R^2$). The kinetic study showed Korsmeyer-Peppas model as best fit model because it showed highest value ($R^2 = 0.9538$). The n value found to 0.54 (0.45 to 0.85) representing non– Fickian mechanism with dual release i.e., diffusion and swelling release (Wu et al., 2019).

6.9. Ex-vivo permeability study

The ex-vivo permeability of AG-BILopt and free AG dispersion was performed on rat intestine. The permeated amount of AG from AG-BILopt was found to be 83.23 ± 3.42 µg/cm² which is significantly higher (p < 0.05) than free AG dispersion (18.76 ± 1.54 µg/cm²). The flux value from AG-BILopt was found to be 1.35 µg/cm²/h, and was 4.49 fold higher than free AG dispersion (0.31 µg/cm²/h). The APC of AG-BILopt and free AG dispersion was also calculated and found to be $1.08 \times 10^{-4}$ cm/min and $2.41 \times 10^{-5}$ cm/min, respectively. The higher permeation was achieved due to nanosized vesicle, high internalization in lipid matrix and also due to presence of nonionic surfactant. The presence of nonionic surfactant has the permeation enhancing capacity, it opens the tight junction of intestine due to more hydrostatic pressure. The surfactant also inhibited the Pgp efflux pump and reduced the reticulo-endothelial uptake. It also lessens the formulation efflux and the bilosomes vesicles having the flexible property due to presence of bile salt (Gaba et al., 2015).

7. Biological study

7.1. Pharmacokinetic study

The pharmacokinetic study of AG-BILopt and free AG-dispersion was performed to evaluate bioavailability of AG. AG concentration vs time profile of AG-BILopt and free AG-dispersion was expressed in Fig. 6. AG-BILopt showed significantly higher (3.86 ± 0.26 µg/ml, P < 0.05) Cmax value than AG-dispersion (1.28 ± 0.11 µg/ml). The higher Cmax of AG-BILopt is due to the smaller size, high encapsulation, high solubility in GIT fluid, high permeability as well as avoids the first-pass metabolism. AUC0-24 and AUC0-$\infty$ value of AG-BILopt treated rats showed 39.92 ± 2.65 (µg.h/ml) and 43.01 ± 2.15 (µg.h/ml). These values are found to be significantly (p < 0.05) higher (4.67 and 4.93 fold) than free AG-dispersion (AUC0-24 8.54 ± 0.98; AUC0-$\infty$ 8.71 ± 1.04). Tmax of AG-BILopt was also found to be high (4 h) as compared to AG-dispersion (2 h). The AUMC0-24 of AG-dispersion is 51.06 ± 1.54 µg.h²/ml whereas AG-BILopt showed 321.01 ± 10.43 µg.h²/ml. The formulation free AG-dispersion and AG-BILopt depicted AUMC0-$\infty$ value of 56.01 ± 3.48 and 421.19 ± 5.64 µg.h²/ml, respectively. The high value of AUC and Tmax is due to slow and prolonged release of AG which helps to absorb maximum drug amount. The elimination rate constant (h⁻¹) of AG-BILopt and free AG-dispersion was found to be 0.1195 and 0.173. Half-life (h) was found to be 3.98 ± 0.25 for free AG-dispersion and 5.79 ± 0.23 AG-BILopt. The overall results indicate AG-BILopt enhances the relative bioavailability of – 4.67 fold as compare to free AG dispersion. The higher bioavailability was found due to the higher uptake of bilosomes by intestinal M–cell of peyer’s patch and also due to increased solubility in the presence of lipid and surfactant (Elnaggar, 2015).

7.2. Evaluation of hypoglycaemic activity

The antihyperglycaemic effect of AG-BILopt and free AG-dispersion were determined by evaluating average fasting blood glucose level (BGL) as depicted in Fig. 7. The normal and diabetic control group rats showed BGL 102 ± 5.9 mg/dl and 205 ± 7.2 mg/dl, respectively. The data showed a reduction of blood glucose level after 1 h administration and the reduction was maintained up to 12 h for AG-BILopt and AG-dispersion showed up to 4 h. The highest blood glucose level reduction was found to be 40.49% (122 ± 4.7 mg/dl in 12 h). The AG-BILopt exhibited remarkable (P < 0.001**, P < 0.0001**) effect than free AG-Dispersion and diabetic control. After 12 h, blood glucose level gradually increases and reached up to 156 mg/dl. The free AG-dispersion showed an increase in BGL after 4 h and reached up to 198 ± 6.7 mg/dl in 24 h. The result of the study indicates AG-BILopt treated animals showed a significant reduction of blood glucose level for a longer time due to enhanced solubility.

7.3. Biochemical evaluation

At the end of the study, the various biochemical parameters (TG, TC, HDL-C, uric acid, urea, serum glutamic–pyruvic transaminase}
Table 5. Comparative estimated biochemical parameters result of different treated groups.

| Groups          | TG (mg/dl) | TC (mg/dl) | HDL-C (mg/dl) | UA (mg/dl) | SGPT (U/L) | SGOT (U/L) |
|-----------------|------------|------------|---------------|------------|------------|------------|
| NC              | 47.76 ± 0.87 | 64.62 ± 0.5 | 50.23 ± 0.6   | 35.72 ± 0.6 | 1.28 ± 0.2 | 28.24 ± 1.03 |
| DC              | 72.2 ± 1.1  | 103.76 ± 1.8 | 28.52 ± 1.6   | 62.76 ± 2.3 | 2.35 ± 0.4 | 51.25 ± 1.43 |
| Free-AG-Dispersion | 57.8 ± 1.2  | 82.39 ± 0.7  | 37.59 ± 1.4   | 51.87 ± 1.5 | 1.87 ± 0.5 | 40.38 ± 1.13 |
| AG-Bilopt       | 49.2 ± 1.4*** | 66.19 ± 0.9*** | 45.39 ± 1.8*** | 39.65 ± 1.4*** | 1.35 ± 0.4*** | 29.28 ± 1.03*** |

***P < 0.0001 as compared to diabetic control group, SGPT: Serum glutamic –pyruvic transaminase, SGOT: Serum glutamic oxaloacetic transaminase, Uric acid:- UA, HDL-C:- High density lipoprotein- Cholesterol, Triglycerides: TG, Total cholesterol: TC.

(SGPT), and serum glutamic oxaloacetic transaminase (SGOT) were evaluated. The biochemical parameters of normal control, diabetic control, treated AG-dispersion and AG-Bilopt was depicted in Table 5. The blood glucose level significantly changed (P < 0.0001) in STZ-induced DM than normal rats. The alteration in lipid profile causes cardiovascular complications associated with type-2 DM (Kaur et al., 2013; Shaveta et al., 2020). The streptozotocin also causes liver toxicity due to alteration in serum glutamic–pyruvic transaminase and serum glutamic oxaloacetic. AG-Bilopt and AG-dispersion treated group exhibited significant (P < 0.0001) reduction in elevated TC and TG as well as the decreased level of glutamic oxaloacetic transaminase was also decreased the elevated level (P < 0.0001) by AG-Bilopt as compared to diabetic control rats. Also, AG-Bilopt significantly decreased the elevated level of total protein in serum as compared to the diabetic control group.

8. Conclusion

In the present study, AG-bilosomes was successfully developed with the objective to enhance the oral efficacy for management of type-2 DM. The prepared AG-BILs have shown nanosize range, high entrapment efficiency, spherical shape, and higher in vitro drug release upto 12 h. The thermal analysis study revealed that AG was encapsulated in lipid matrix. The intestinal permeation study revealed that high amount of AG permeated (P < 0.05) than free AG-dispersion. Pharmacokinetic study showed AG-Bilopt enhances the systemic bioavailability and residence time than free AG-dispersion. The pharmacodynamic study also revealed a significant (P < 0.05) enhancement in the hypoglycemic activity and biochemical parameters than free AG. Our findings concluded that oral AG-BILs was found to be better treatment alternative for diabetes with improved therapeutic efficacy.

Declaration of Competing Interest

The authors declare that they have no competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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