Biosynthesis, characterization and anti-dengue vector activity of silver nanoparticles prepared from *Azadirachta indica* and *Citrullus colocynthis*

Shafqat Rasool, Muhammad Akram Raza, Farkhanda Manzoor, Zakia Kanwal, Saira Riaz, Muhammad Javaid Iqbal and Shahzad Naseem

Article citation details
*R. Soc. open sci.* 7: 200540.
http://dx.doi.org/10.1098/rsos.200540

Review timeline
Original submission: 1 May 2020
1st revised submission: 4 July 2020
2nd revised submission: 6 August 2020
Final acceptance: 13 August 2020

Note: Reports are unedited and appear as submitted by the referee. The review history appears in chronological order.

Review History

RSOS-200540.R0 (Original submission)

Review form: Reviewer 1 (Mudassir Iqbal)

Is the manuscript scientifically sound in its present form?
Yes

Are the interpretations and conclusions justified by the results?
Yes

Is the language acceptable?
Yes

Do you have any ethical concerns with this paper?
No

Have you any concerns about statistical analyses in this paper?
No
Recommendation?
Accept with minor revision (please list in comments)

Comments to the Author(s)
The article describes the synthesis of Ag nanoparticle using Azadirachta indica and Citrullus colocynthis extracts. The work done is useful and is in line with the current scientific demands. The are also some issue to be addressed before publishing this work.

- There are several grammatical mistakes for instance I am highlighting some of these:
  i) page 2 line 17 which have famous is incorrect it should be which are famous
  ii) page 2 line 25 “vector for many deceases like” it should be vector for many diseases like
  iii) page 2 line 25 please correct “fever.Dengue”
  iv) page 2 line 35 “AgNPs were manufactured” were synthesized
  v) page 2 line 35 “the mixture was let to cool down to room temperature” should be the mixture was cooled to room temperature

The authors should thoroughly check the whole manuscript and English should be improved as well as all the editorial mistakes should be rectified.

- Page 6 line 10, authors mentioned the particle size, I guess it should be crystallite size?
- Authors have not mapped the particle size in SEM images. The particle size should be measured in SEM image and it should be visible in the picture.
- The authors have mentioned that the chosen plants have known biological activity. However, they do not show any activity against A. aegypti. Secondly, if the extracts are not active against A. aegypti why there is a large difference in activity of AgNPs prepared from two different extracts as LC50 at 1.25 ppm for A. indica and LC50 at 0.3 ppm for C. colocynthis is observed. The later is 4 times more active than the former one. i does not seem that activity is only due to Ag NPs in that case it should be similar or close to each other.
- Figures 1,2 and 5 are unnecessary and should be deleted.
- the sentence in conclusion section “Both types of NPs were found pure metallic silver in nature having FCC crystalline structure and of spherical in shapes with diameters in the range of 25±5 nm, synthesized by different biomolecules of extracts as determined by different characterization techniques including SEM, UV- Vis, XRD and FTIR” is too long and ununderstandable. It should be refabricated to two sentences.

After addressing these issues the article can be published in RSOS.

Review form: Reviewer 2

Is the manuscript scientifically sound in its present form?
No

Are the interpretations and conclusions justified by the results?
No

Is the language acceptable?
No

Do you have any ethical concerns with this paper?
No

Have you any concerns about statistical analyses in this paper?
Yes

Recommendation?
Major revision is needed (please make suggestions in comments)
Comments to the Author(s)
The MS presents work which has been published with various other plants. The work carried out is significant but need major revision.

MS needs enormous revision in scientific reporting, language and grammar. The scientific names are wrongly spelled at some places.

Each section of the MS needs major revision. The comments have been marked on the MS (Appendix A). Authors are advised to go through each and every comment; and rectify accordingly.

Decision letter (RSOS-200540.R0)

We hope you are keeping well at this difficult and unusual time. We continue to value your support of the journal in these challenging circumstances. If Royal Society Open Science can assist you at all, please don't hesitate to let us know at the email address below.

Dear Dr Raza:

Title: Biosynthesis, Characterization and Anti-dengue activity of Silver Nanoparticles prepared by Azadirachta indica and Citrullus colocynthis
Manuscript ID: RSOS-200540

Thank you for your submission to Royal Society Open Science. The chemistry content of Royal Society Open Science is published in collaboration with the Royal Society of Chemistry.

The editor assigned to your manuscript has now received comments from reviewers. We would like you to revise your paper in accordance with the referee and Subject Editor suggestions which can be found below (not including confidential reports to the Editor). Please note this decision does not guarantee eventual acceptance.

Please submit your revised paper before 05-Jul-2020. Please note that the revision deadline will expire at 00.00am on this date. If we do not hear from you within this time then it will be assumed that the paper has been withdrawn. In exceptional circumstances, extensions may be possible if agreed with the Editorial Office in advance. We do not allow multiple rounds of revision so we urge you to make every effort to fully address all of the comments at this stage. If deemed necessary by the Editors, your manuscript will be sent back to one or more of the original reviewers for assessment. If the original reviewers are not available we may invite new reviewers.

To revise your manuscript, log into http://mc.manuscriptcentral.com/rsos and enter your Author Centre, where you will find your manuscript title listed under "Manuscripts with Decisions." Under "Actions," click on "Create a Revision." Your manuscript number has been appended to denote a revision. Revise your manuscript and upload a new version through your Author Centre.

When submitting your revised manuscript, you must respond to the comments made by the referees and upload a file "Response to Referees" in "Section 6 - File Upload". Please use this to document how you have responded to the comments, and the adjustments you have made. In order to expedite the processing of the revised manuscript, please be as specific as possible in your response.
Once again, thank you for submitting your manuscript to Royal Society Open Science and I look forward to receiving your revision. If you have any questions at all, please do not hesitate to get in touch.

Yours sincerely,
Dr Laura Smith
Publishing Editor, Journals

Royal Society of Chemistry
Thomas Graham House
Science Park, Milton Road
Cambridge, CB4 0WF
Royal Society Open Science - Chemistry Editorial Office

On behalf of the Subject Editor Professor Anthony Stace and the Associate Editor Dr Ya-Wen Wang.

******************************************************************************

RSC Associate Editor:
Comments to the Author:
(There are no comments.)

RSC Subject Editor:
Comments to the Author:
(There are no comments.)

******************************************************************************

Reviewers' Comments to Author:
Reviewer: 1

Comments to the Author(s)
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After addressing these issues the article can be published in RSOS.

Reviewer: 2

Comments to the Author(s)
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Each section of the MS needs major revision. The comments have been marked on the MS. Authors are advised to go through each and every comment; and rectify accordingly.

Author’s Response to Decision Letter for (RSOS-200540.R0)
See Appendix B.

RSOS-200540.R1 (Revision)

Review form: Reviewer 1 (Mudassir Iqbal)

Is the manuscript scientifically sound in its present form?
Yes

Are the interpretations and conclusions justified by the results?
Yes

Is the language acceptable?
Yes

Do you have any ethical concerns with this paper?
No

Have you any concerns about statistical analyses in this paper?
No

Recommendation?
Accept as is

Comments to the Author(s)
Nil
Review form: Reviewer 2

Is the manuscript scientifically sound in its present form?
Yes

Are the interpretations and conclusions justified by the results?
Yes

Is the language acceptable?
Yes

Do you have any ethical concerns with this paper?
No

Have you any concerns about statistical analyses in this paper?
No

Recommendation?
Accept with minor revision (please list in comments)

Comments to the Author(s)
I appreciate the efforts taken in revision of the manuscript. A few errors are yet to be addressed. The errors have been incorporated in the MS. Authors are requested to go through the MS (Appendix C) and address them.

Decision letter (RSOS-200540.R1)

We hope you are keeping well at this difficult and unusual time. We continue to value your support of the journal in these challenging circumstances. If Royal Society Open Science can assist you at all, please don't hesitate to let us know at the email address below.

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Title: Biosynthesis, Characterization and Anti-dengue vector activity of Silver Nanoparticles prepared by Azadirachta indica and Citrullus colocynthis
Manuscript ID: RSOS-200540.R1

Thank you for submitting the above manuscript to Royal Society Open Science. On behalf of the Editors and the Royal Society of Chemistry, I am pleased to inform you that your manuscript will be accepted for publication in Royal Society Open Science subject to minor revision in accordance with the referee suggestions. Please find the reviewers' comments at the end of this email.

The reviewers and handling editors have recommended publication, but also suggest some minor revisions to your manuscript. Therefore, I invite you to respond to the comments and revise your manuscript.

Because the schedule for publication is very tight, it is a condition of publication that you submit the revised version of your manuscript before 13-Aug-2020. Please note that the revision deadline will expire at 00.00am on this date. If you do not think you will be able to meet this date please let me know immediately.
To revise your manuscript, log into https://mc.manuscriptcentral.com/rsos and enter your Author Centre, where you will find your manuscript title listed under "Manuscripts with Decisions". Under "Actions," click on "Create a Revision." You will be unable to make your revisions on the originally submitted version of the manuscript. Instead, revise your manuscript and upload a new version through your Author Centre.

When submitting your revised manuscript, you will be able to respond to the comments made by the referees and upload a file "Response to Referees" in "Section 6 - File Upload". You can use this to document any changes you make to the original manuscript. In order to expedite the processing of the revised manuscript, please be as specific as possible in your response to the referees.

When uploading your revised files please make sure that you have:

1) A text file of the manuscript (txt, txt, rtf, docx or doc), references, tables (including captions) and figure captions. Do not upload a PDF as your "Main Document".
2) A separate electronic file of each figure (EPS or print-quality PDF preferred (either format should be produced directly from original creation package), or original software format)
3) Included a 100 word media summary of your paper when requested at submission. Please ensure you have entered correct contact details (email, institution and telephone) in your user account
4) Included the raw data to support the claims made in your paper. You can either include your data as electronic supplementary material or upload to a repository and include the relevant doi within your manuscript
5) All supplementary materials accompanying an accepted article will be treated as in their final form. Note that the Royal Society will neither edit nor typeset supplementary material and it will be hosted as provided. Please ensure that the supplementary material includes the paper details where possible (authors, article title, journal name).

Supplementary files will be published alongside the paper on the journal website and posted on the online figshare repository (https://figshare.com). The heading and legend provided for each supplementary file during the submission process will be used to create the figshare page, so please ensure these are accurate and informative so that your files can be found in searches. Files on figshare will be made available approximately one week before the accompanying article so that the supplementary material can be attributed a unique DOI.

Once again, thank you for submitting your manuscript to Royal Society Open Science. The chemistry content of Royal Society Open Science is published in collaboration with the Royal Society of Chemistry. I look forward to receiving your revision. If you have any questions at all, please do not hesitate to get in touch.

Kind regards,
Dr Laura Smith
Publishing Editor, Journals
Royal Society of Chemistry
Thomas Graham House
Science Park, Milton Road
Cambridge, CB4 0WF
Royal Society Open Science - Chemistry Editorial Office

On behalf of the Subject Editor Professor Anthony Stace and the Associate Editor Dr Ya-Wen Wang.
Author's Response to Decision Letter for (RSOS-200540.R1)

See Appendix D.

Decision letter (RSOS-200540.R2)

We hope you are keeping well at this difficult and unusual time. We continue to value your support of the journal in these challenging circumstances. If Royal Society Open Science can assist you at all, please don't hesitate to let us know at the email address below.

Dear Dr Raza:

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Manuscript ID: RSOS-200540.R2

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The comments of the reviewer(s) who reviewed your manuscript are included at the end of this email.
Thank you for your fine contribution. On behalf of the Editors of Royal Society Open Science and the Royal Society of Chemistry, I look forward to your continued contributions to the Journal.

Yours sincerely,
Dr Laura Smith
Publishing Editor, Journals

Royal Society of Chemistry
Thomas Graham House
Science Park, Milton Road
Cambridge, CB4 0WF
Royal Society Open Science - Chemistry Editorial Office

On behalf of the Subject Editor Professor Anthony Stace and the Associate Editor Dr Ya-Wen Wang.

********

RSC Associate Editor
Comments to the Author:
(There are no comments.)

********

Reviewer(s)' Comments to Author:
Appendix A

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Biosynthesis, Characterization and Anti-dengue activity of Silver Nanoparticles prepared by Azadirachta indica and Citrullus colocynthis

| Journal:          | Royal Society Open Science |
|-------------------|----------------------------|
| Manuscript ID     | RSOS-200540                |
| Article Type:     | Research                   |
| Date Submitted by the Author: | 01-May-2020               |
| Complete List of Authors: | Rasool, Shafqat; University of the Punjab, Quid-e-Azam Campus, Lahore-54590, Pakistan, Centre of Excellence in Solid State Physics Raza, Muhammad Akram; University of the Punjab, Centre of Excellence in Solid State Physics Manzoor, Farkhanda ; Lahore College for Women University, Department of Zoology Kanwal, Zakia; Lahore College for Women University, Jail Road, Lahore-54000,, Department of Zoology Riaz, Saira; University of the Punjab, Centre of Excellence in Solid State Physics Iqbal, Muhammad Javaid; University of the Punjab, Centre of Excellence in Solid State Physics Naseem, Shahzad ; University of the Punjab, Centre of Excellence in Solid State Physics |
| Subject:          | Green chemistry < CHEMISTRY, Nanotechnology < CHEMISTRY, biophysics < CROSS-DISCIPLINARY SCIENCES |
| Keywords:         | anti-dengue activity, silver nanoparticles, green synthesis, Citrullus colocynthis, Azadirachta indica |
| Subject Category: | Chemistry                   |

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Author-supplied statements

Relevant information will appear here if provided.

Ethics

Does your article include research that required ethical approval or permits?:
This article does not present research with ethical considerations

Statement (if applicable):
CUST_IF_YES_ETHICS :No data available.

Data

It is a condition of publication that data, code and materials supporting your paper are made publicly available. Does your paper present new data?:
Yes

Statement (if applicable):
Data Accessibility statement:

Our data are deposited at Dryad:
https://datadryad.org/stash/dataset/doi:10.5061/dryad.z612jm68k [48], with following reviewer URL;

Dryad Reviewer URL:
https://datadryad.org/stash/share/DksW82FCdqlSwPql2aYhDi4C9QUE74hEBPTmrJ3trHw

Conflict of interest

I/We declare we have no competing interests

Statement (if applicable):
CUST_STATE_CONFLICT :No data available.

Authors' contributions

This paper has multiple authors and our individual contributions were as below

Statement (if applicable):
Authors' Contributions:

1). SR and MAR performed the synthesis experiments, analysed the data and wrote the manuscript
2). ZK, SR and MAR designed the experiments and conducted the larvicidal bioassays
3). SR and SN conducted the characterization measurements and helped in interpretation of data
4). FM and ZK provided the larvae facility and helped in larvicidal activity experiments
5). SR and SN conducted the characterization measurements and helped in interpretation of data
6). MJR and SR conducted the FTIR and SEM characterization and helped in data analysis
Biosynthesis, Characterization and Anti-dengue activity of Silver Nanoparticles prepared by *Azadirachta indica* and *Citrullus colocynthis*

Shafqat Rasool\(^1\), Muhammad Akram Raza\(^*\),\(^1\) Farkhanda Manzoor\(^2\), Zakia kanwal\(^2\), Saira Riaz\(^1\), Muhammad Javaid Iqbal\(^1\), Shahzad Naseem\(^1\)

\(^1\)Centre of Excellence in Solid State Physics, University of the Punjab, Quid-e-Azam Campus, Lahore 54590, Pakistan

\(^2\)Department of Zoology, Lahore College for Women University, Jail Road, Lahore 54000, Pakistan

**Keywords:** anti-dengue activity, silver nanoparticles, green synthesis, *Citrullus colocynthis*, *Azadirachta indica*

**Abstract**

Green synthesis of nanomaterials using different bio-products can minimize the negative side effects of nanotechnology. Here, we report biosynthesis of silver nanoparticles (AgNPs) using aqueous extracts of (i) *Azadirachta indica* leaves and (ii) *Citrullus colocynthis* fruit and their larvicidal activity against *Aedes Aegypti*. The prepared AgNPs were characterized by Fourier transforms infrared (FTIR), x-ray diffraction (XRD), uv-vis spectroscopy and scanning electron microscopy (SEM) for chemical, structural, optical and morphological analysis. Pure silver nature of colloidal NPs was indicated by occurrence of uv-vis absorption peaks in the range of 420-430 nm whereas XRD pattern confirmed the face centred cubic (FCC) structure of NPs. SEM examination revealed the spherical morphology of AgNPs with size 25±5 nm while characteristic peaks appearing in FTIR analysis indicated the attachment of different biomolecules on AgNPs. The anti-dengue activity of synthesized AgNPs at different concentrations (1ppm to 30 ppm) and as-prepared extracts against *Aedes aegypti* larvae were examined for 24 h. A concentration dependent larvicidal potential of both types of AgNPs was observed, however, *C. colocynthis* mediated AgNPs (LC\(_{50}\) at 0.3 ppm) were found more effective at lower concentration than *A. indica* mediated AgNPs (LC\(_{50}\) at 1.25 ppm). However, extracts did not exhibit any significant larvicidal activity.

1. Introduction

Nanotechnology, owing to unique, extraordinary and incredible properties of materials at nano-scale, has revolutionized almost every field of science and technology. Especially, in arena of medical science and medicine, it is considered as next logical step and the future medicine [1, 2]. However, the potential risks such as harmful side effects on human and environment has made nanotechnology a double-edged sword [3,4].

Natural products based synthesis of nanoparticles can help to reduce the hazards of nanotechnology and thus can boost its applications. Biosynthesis, based on green chemistry principles, is proactive approach and has numerous advantages over conventional chemical and physical methods such as cost-effectiveness, environment friendly, simple and safe, no use of hazardous chemicals as reducing agents, less wastage of materials, minimum energy usage, safer disposal and recycling [5, 6]. Studies are also reported on the preparation of various types of nanostructures by using natural products such as vitamins, biodegradable polymers, enzymes, polysaccharides and different microorganism including algae, fungi, bacteria and viruses. Nevertheless, plants extract mediated synthesis of nanomaterials is consider advantageous due to stability of...
nanostructures, cost efficient, availability of plants, lower contamination risks, require little maintenance, and simple to scale up as plants are nature’s ‘chemical factories’ [6,7]. Silver Nanoparticles are famous for their anti-bacterial, anti-dengue and anti-fungal activate gNPs are also used in textile industry, food packing, cosmetics, paints and detergents owing to their anti-microbial activity. Preparation of AgNPs can be carried out by using extract of different parts of plant such as roots, stem, leaves or fruit. In the synthesis of AgNPs by plants, polyphenols play the main role in degradation of different organics compounds. These plant extracts usually play the dual role in the synthesis process as reducing and capping agent. During green synthesis, the coatings of various biomolecules on surfaces of AgNPs not only improve their stability but also enhance their biocompatibility by reducing the toxicity risks [8, 9].

In this study, we choose Azedarachta Indica (A. Indica) and Citrullus Colocynthis (C. Colocynthis) for the synthesis of AgNPs because both of these plants are famous for good medicinal properties and exhibit effective biological activities. A. Indica plant belongs to family of Meliaceae and has enormous uses in medical field from ancient times. It is familiar as anti-bacterial, anti-fungal and anti-microbial plant as it has quercetin, β-sitosterol and nimbinin in their leaves and azadirachtin in its seeds. The A. Indica plant extract has been reported to increases anti-microbial activity of extract mediated nanoparticles [10, 11]. C. Colocynthis (also known as bitter apple) is member of Cucurbitaceae family. It contains many biomolecules including alkaloids, glycosides, flavonoids, and fatty acids which have famous for important biological activities such as antimicrobial, antioxidants, cytotoxic, anti-diabetic, antilipidemic, and insecticide. That’s why it is considered one of the best medicinal plants and is used in different biomedical applications including anti-microbial, anticancer activities [12-14].

According to World Health Organization (WHO), mosquitoes can be listed as one of the deadliest organisms because millions of humans die due to different diseases caused by them. Mosquitoes are the potential vector for many deceases like dengue, malaria, west nile, chikungunya, zika and yellow fever.Dengue incidence has increased 30-fold worldwide in the last 30 years which is caused by dengue mosquito larvae: Aedes aegypti (A. aegypti) [15]. Treatment of these diseases is a challenge. To control the rapidly increasing mosquito-borne risks, researchers are working on different strategies. Since mosquitoes breed in water and at larval stage, it is easy to target them there. However, inhibiting larvae in water by conventional ways using pesticides can increase the toxic risks for environment and humans. Thus natural pesticides such as plant extracts can be simple and side effect free promising methodologies. The use of green chemistry based nanomaterials can further enhance the effectiveness and efficiency of such approaches [16-19].

In this work, A. Indica and C. Colocynthis mediated AgNPs were manufactured, characterized and tested for their anti-dengue activity and found that these AgNPs can potentially be considered good anti-larvicidal agents.

2. Materials and Methods

2.1. Material and chemicals

Fresh leaves of A. indica were collected from the Botanical garden of University of the Punjab, Lahore, Pakistan and C. colocynthis fruit was obtained from local market of Lahore, Punjab, Pakistan. Both products were identified by the experts of Department of Botany, University of the Punjab, Lahore. Late fourth instar larvae of A. aegypti were collected from local pond and identified by the expert of the Department of Zoology, Lahore College for Women University, Lahore, Pakistan. Silver nitrate (AgNO₃) was of research grade and obtained from Meck (Germany). To prepare synthesis solutions, aqueous extracts and for all other purposes deionized (DI) water was used throughout the experiment.

2.2. Preparation of A. indica and C. colocynthis extracts

The aqueous extract of A. indica leaves was prepared following the method reported by Pragyan et al [20] with some modifications. To remove any dust and contaminations, first fresh leaves were rinsed carefully with running tap water then washed twice with DI water and let them dry in air. 20g of these leaves were taken and converted into very small pieces by cutting them transferred to conical flask of 250ml to boil for 10 min in 100ml deionized water. After boiling, the mixture was let to cool down to room temperature and finally filtered with Whatman No. 1 filter paper before storing at 4 °C for next step. Different steps of the preparation of A. indica leaves aqueous extract are shown in Figure 1. One can notice that greenish color of mixture solution (figure 1c, before boiling) changed to yellowish light brown color in final stage (figure 1d).
The aqueous extract of *C. colocynthis* fruit was prepared by the technique mentioned elsewhere [21] with few changes and whole process of extract preparation is demonstrated in figure 2. Briefly, a greenish white *C. colocynthis* fruit (as shown in figure 2a) was cleaned by washing many times with fresh tap water to get rid of any debris or any other contaminations, then rinsed two times with DI water before cutting into small pieces. 50g pieces of *C. colocynthis* were soaked in 200ml of deionized water for 72h. Finally mixture was filtered 2 times with Whatman No.1 filter paper to get greenish yellow color of the extract (figure 2c) and kept at 4ºC for further use.

### 2.3. Synthesis of extract mediated silver nanoparticles

To prepare AgNPs using aqueous extracts, first precursor stock solution (100mM) was made by dissolving 1.69 g of silver nitrate (AgNO₃) into 100ml of DI water.

*A. indica* extract mediated AgNPs were made by the protocol of Aparajita et al. [22] with some amendments. We, in our case, used three different amounts of as-prepared extract (250µl, 500µl and 1000µl) for the same amount 20ml of silver nitrate precursor solution (1mM) to obtain the optimized concentration. In figure 3 (upper panel), different phases of AgNPs preparation using the *A. indica* extract are presented. First of all colorless precursor solution of AgNO₃ was heated to 70°C with gentle magnetic stirring of 200 rpm (figure 3a). Then *A. indica* extract was added dropwise into the solution and the mixture was kept for heating at hot plate at 70°C under continuous stirring for 10 min (figure 3b). The color of the solution changed with time from transparent to light yellow to yellowish brown indicating the formation of AgNPs (figure 3c). At the end, the solution was let to cool down to room temperature and kept at -4°C for further use. Depending upon the amount of the extract the color of final stage sample (colloidal AgNPs) was found to change as can be noticed in lower panel of figure 3 (lower panel), from yellowish brown to reddish brown to reddish dark brown for 250 µl, 500 µl and 1000µl extract mediated AgNPs respectively. However, only AgNPs prepared by using the 500 µl of extract was used in larvicidal activity and other characterization.

For the synthesis of AgNPs using aqueous extract of *C. colocynthis*, approach described by Shawkey et al. [23] was followed with some alterations. Briefly, first 20ml of 5mM silver nitrate solution was mixed with a defined amount of *C. colocynthis* extract then the mixture was heated to the boiling temperature (about 95°C) under constant stirring of 200 rpm for 20 min. Finally, solution was cooled down to room temperature and store at -4°C. Color changes during the synthesis process indicate different phases of the synthesis and are presented in figure 4 (upper panel). We synthesized AgNPs using three amounts of as-prepared extract of *C. colocynthis* viz., 1ml, 2.5 ml and 4 ml following aforementioned method to achieve the optimized concentration. All three colloidal AgNPs samples mediated by different amounts of extract with precursor and extract samples are shown in figure 4 (lower panel). Nevertheless, only 2.5 ml extract mediated AgNPs were further used for larvicidal activity and other characterization.

### 2.4. Characterization of synthesized silver nanoparticles

The above prepared AgNPs were characterized by different techniques. Optical properties of all colloidal samples were examined by measuring the absorbance spectra with the ultraviolet–visible (UV-Vis) spectroscopy (Shimadzu, UV-1800, Japan) in the range of 300-700nm wavelength. The size, shape and elemental composition investigations of synthesized nanoparticles were conducted by scanning electron microscopy and energy-dispersive X-ray (Nova NanoSEM 450, USA). For SEM analysis, samples were prepared by using the drop casting methods to achieve sufficient AgNPs amount on the clean glass substrate. To avoid any probable charging effects, a very thin gold coating was deposited on the samples before SEM conducting. The structural nature of prepared NPs was examined by X-ray diffractometer (JSX 3201M, Jeol, Japan). For XRD, again, samples were prepared by developing a thick enough film of each colloidal sample by drop casting methods on the glass substrate. To study the presence of different probable functional group on the surface of AgNPs, fourier transforms infrared spectroscopy (IRTracer-100, shimadzu, Japan) was conducted in the range of 500-3600 cm⁻¹.

### 3. Larvicidal activity tests

The larvicidal potential of both types of green AgNPs and their respective extracts at different concentrations against dengue larvae were studies according to the standard protocol recommended by WHO [24]. 25 fourth
instar *A. aegypti* larvae collected from local pond were transferred into different beakers each containing 200 ml deionized water. Five different concentrations of AgNPs were prepared and tested in each case; for *A. Indica* mediated AgNPs following concentrations were used: 1.25 ppm, 2.5 ppm, 5 ppm, 10 ppm and 20 ppm and for *C. colocynthis* mediated AgNPs concentrations used were: 0.3 ppm, 0.6 ppm, 1.25 ppm, 2.5 ppm and 5 ppm. The test concentrations of both types of AgNPs were selected on the basis of pilot experiments in which LC₅₀ concentrations were found. In each beaker required amount of AgNPs were added to achieve to the predetermined concentration. In the case of extracts, three concentrations (1 ml, 2 ml and 3 ml) of each type of extracts were chosen while only DI water was used as control. The experiment was conducted in triplicate under laboratory conditions (at 26 °C). All the beakers were covered with perforated aluminum foil for air circulation. During exposure experiment, larvae were provided no food. After 24 h of treatment period, the counting of alive and dead larvae was made to calculate percent mortality. Larvae which showed no motility after being disturbed with a needle were termed as dead. The data are shown as mean value and standard error of mean (mean+ s.e.); and Microsoft Excel program was used to make the graphs. The percentage mortality was determined by using formula given below

\[
\text{% Mortality} = \frac{\text{Number of dead larvae}}{\text{total number of larvae exposed}} \times 100
\]

4. Results and discussion

4.1. Synthesis mechanism of extract mediated AgNPs

Figure 5 illustrates schematically the potential formation mechanism of AgNPs using the aqueous extracts of different biomaterials. First of all, the precursor (AgNO₃) is dissolved into the solvent (DI water) that provides Ag-ions (Ag⁺) into solution without producing any color. Addition of extract induces the reduction of these Ag-ions to free Ag-atoms (Ag⁰) by gaining the electrons. This reduction can be termed as bio-reduction or plant-assisted reduction because it is caused by different biomolecules such as protein and phytochemicals present in the extract which serve as reducing agent [25]. This reduction reaction can be noticed by the color change of the solution which may be enhanced by heating. The color of the reaction solution may change from transparent to light yellow to yellowish brown and finally dark brown depending upon the type of the extract used (as shown in figures 3 and 4). This time dependent color change during the synthesis, actually, indicates the different phases of nucleation and growth process occurring in the reactions [26]. The free silver atoms (Ag⁰) accumulate due to van der Waals interactions and Brownian motion to form Ag-nuclei (nucleation process). The growth process of these nuclei into AgNPs can occur in different ways including coalescence of these nuclei and Ostwald ripening of smaller NPs into bigger ones. However, both nucleation and growth process may occur simultaneously in reaction during the formation of AgNPs [27]. The biomolecules of the extracts also served as capping agents because no additional stabilizing agent was used. The final size of the synthesized AgNPs can be controlled by varying the concentrations of extracts and precursor solution [28].

4.2. UV-Vis spectroscopy analysis

UV-Vis spectroscopy is very common and effective way to study the formation of colloidal nanoparticles by their optical responses. In Metallic NPs such as AgNPs, the free movement of electron between the valance band and conduction bands is possible due to very narrow gap between them. These electrons on the surface of AgNPs produce the surface plasmon resonance (SPR) which is actually resonant oscillation of conduction electrons in response of the incident light. Due to the SPR absorption nanoparticles in colloidal form produce different colors depending upon various factors e.g., size, shape and surrounding medium [25].

To confirm purity and to study the optical response of colloidal AgNPs, UV-Vis spectroscopy was conducted and obtained absorption spectra are displayed in figure 6. The characteristics surface plasmon resonance (SPR) peaks of AgNPs using three different amounts of *A. Indica* extract 250 µl, 500 µl and 1000 µl are shown in left panel (figure 6a). All three peaks were in the wavelength range of 412 nm to 416 nm indicating the pure silver nature of the particles. A slight red shift can be noticed with increasing concentration of the extract in the reaction solution. A similar trend of peak shifting towards the higher wavelength values (416 nm, 429 nm, 431 nm) by enhancing the amount of extract (1 ml, 2.5 ml and 4 ml) was also observed in the synthesis of *C. colocynthis* mediated AgNPs as illustrated in right panel (figure 6b). This red shift might be due to the change in size of the AgNPs because, in green synthesis, extracts served as the
4.3. Scanning electron microscope (SEM) analysis

SEM is the most prominent characterization technique to analyze shape, surface morphology, size distribution and elemental composition of nanomaterials. In order to examine the morphology of AgNPs prepared by both extracts, SEM analysis was conducted and the obtained micrographs with histograms of particle size distribution are presented in figure 7.

In case of AgNPs mediated from A. Indica extract, small spherical shaped silver particles like bright tiny spots can be seen on the bigger bio-particles of the extract. At some places agglomeration of few smaller particles into small clusters can also be noticed (figure 7a). There could be many reasons for agglomeration of these nanoparticles because these particles have very high surface energy and to reduce energy particles agglomerate and form larger sized particles [32]. The size distribution range of these spherical AgNPs was found to be from 12 nm to 24 nm (average particle size ~ 17±4 nm).

In the lower panel, SEM micrograph (figure 9c) reveals the size and morphology of AgNPs mediated from C. colocynthis extract. A nice random distribution of NPs over the bigger extract particles can be visualized. No cluster formation due to agglomerate of NPs was noticed. Again most of the NPs were found of spherical shapes with average diameter of 26±5 nm (figure 9d).

To confirm the existence of elemental silver in the synthesized particles, Energy-dispersive X-ray (EDX) was conducted. EDX technique determines the elemental composition analysis on the basis of energy values of characteristics x-rays which are unique for every element. The obtained EDX results are presented in figure 8, in upper panel for A. indica extract mediated AgNPs while in lower panel for C. colocynthis extract mediated particles. Insets show the table of elements in each case.

In both cases, we can notice strong silver signal along with some other signals. The intense optical absorption peak at around 3 KeV is considered a typical characteristic absorption band for metallic silver due to surface plasmon resonance [28]. Thus EDX established the presence of elemental silver in both types of extract mediated NPs by the characteristic SPR absorption peaks which supported the UV-vis results. The other signals such as C, O, Na, Mg, Au, Cl, and K in the case of by A. indica AgNPs and C, O, Na, Ca, Mg, Si, Au, Cl, Pd, and K (figure 8a) for the C. colocynthis AgNPs sample (figure 8b) can also be observed. The occurrence of weak gold (Au) signal can be attributed to thin gold coatings on the sample to avoid any charging. The Si peak may be due to presence of glass substrate. The appearance of other signals especially intense O and C peaks indicate the presence of other metabolites on the AgNPs surface due to the aqueous extract (A. indica leaves and C. colocynthis fruit). These metabolites play a vital role of capping and stabilizing agents in the biosynthesis of the AgNPs because they provide the stability to AgNPs by surrounding and developing a thin capping layer of organic molecules. This advantage of green synthesis approach not only reduce the cost but also minimize the toxicity caused by hazardous chemicals used for reducing and capping purposes in chemical methods [28,33,34].

4.4. X-Ray Diffraction (XRD) analysis

For the structural analysis of materials, X-ray diffraction (XRD) is an eminent analytical technique. It is widely used as primary analytical tool to investigate material purity and different structural parameters such as crystal structure, crystallite size, lattice parameter, crystal phase identification, and various crystal defects [25]. In figure 9, indexed XRD patterns of both types of extract mediated AgNPs are exhibited.

One can notice six prominent peaks in the diffractogram of A. Indica mediated AgNPs peaks (figure 9a). The peaks appearing at 2θ values of 38.5º, 46.5º, 64.7º and 77.1º can be assigned to diffraction planes (hkl values) of (111), (200), (220) and (311) respectively, according to COD ID No. 9013052 [35]. These distinct reflection planes confirmed the silver metallic nature with face centered cubic (FCC) crystalline structure of synthesized AgNPs. Moreover, two other unidentified intense peaks occurring at 2θ values of 28.1º and 32.4º, showing higher degree of crystallinity, can also be noted (labeled by stars). These peaks indicate the presence of some crystalline moieties or organic compounds deposited on surface of AgNPs from aqueous extract of A. indica leaves. Many other studies of AgNPs prepared by green synthesis using the plant extract have also mentioned the appearance of such additional peaks in their XRD spectra which are believed to occur due to crystalline impurities of extracts [36, 37].
In XRD pattern of AgNPs prepared by C. colocynthis as shown figure 9b, four prominent distinct diffraction peaks observed at 2θ value of 37.9° (111), 46.2° (200), 67.5° (220) and 77.1° (311) indicate the FCC crystal structure of resultant nanoparticle, according to COD ID No. 9013046 [38]. Again, some extra intense peaks at 27.8° and 32.2° were observed (star peaks) which may be attributed to existence of crystalline nature of biomolecules on the AgNPs surface due to C. colocynthis extract [39].

To determine the crystallite size (D) of both type of extract mediated AgNPs, the width of prominent Bragg’s reflection (111) in each case was utilized in the Debye–Scherer formula [26].

\[
D = \frac{k \lambda}{\beta \cos \theta}
\]

In this equation (4.1), \(\lambda\), k, \(\beta_{FWHM}\) and \(\theta\) are x-ray wavelength (1.54 Å), shape factor with value of 0.9, peak width (full width half maximum- FWHM) and Bragg diffraction angle respectively. The average particle size determined by XRD for AgNPs prepared using extracts of A. indica and C. colocynthis were found to be 11±1 nm and 15±1 nm respectively (table 1) which has fair agreement with particle size measurements obtained by SEM.

| Table 1 Calculated values of crystallite size and lattice parameter for both types of AgNPs. |
|---------------------------------------------------------------|
| Sample | Peak position (2θ) | Diffraction plane (hkl) | FWHM (rad) | Crystallite size, D (nm) | Lattice parameter, a (nm) |
|-----------------|-----------------|------------------|-------------|-----------------|-----------------|
| A. indica extract mediated AgNPs | 38.5° | (111) | 0.0128 | 11±1 | 0.403 |
| C. colocynthis extract mediated AgNPs | 37.9° | (111) | 0.0095 | 15±1 | 0.409 |

4.5. Fourier-Transform Infrared Spectroscopy (FTIR) Analysis

FTIR spectroscopy is a very useful and powerful technique to identify different functional groups and chemical bonds in any sample. FTIR characterizes the chemical composition of the material by interaction of infrared radiation with a surface of the sample. The infrared radiations are absorbed by the sample at specific frequency ranges depending upon the type of chemical functional groups and bonds. The measurement of these frequencies and intensities help to study the chemistry of the sample.

In this work, to study the presence of different biomolecules and functional groups on the surface of AgNPs, in both cases, which provided the required bio-reduction and efficient stabilization during the synthesis process, FTIR measurement were carried out and obtained spectra are presented in figure 10.

FTIR spectrum of AgNPs synthesized from A. Indica leaves is shown in figure 10a where various absorption bands at different specific frequency indicate a composite nature. The prominent peaks occurred at the frequency ranges of 3265, 2919, 2849, 2363, 1734, 1616, 1244, 1161, 1097 and 1027 cm\(^{-1}\) as can be noticed from the figure 10a. A broad band appearing in the range of 3265 cm\(^{-1}\) can be attributed to the stretching vibrations of O-H bond indicating the presence of polyphenols. The peaks occurring at value range of 2919 cm\(^{-1}\) and 2849 cm\(^{-1}\) can be assigned to C=C aromatic and C-H alkaline (asymmetric stretching of C-H bonds) groups [40]. The possible presence of C=N and C=C triple bonds can be ascribed by peaks of 2363 cm\(^{-1}\) because band appearing in the range of 2200-2400 cm\(^{-1}\) indicate the C-N and C-C triples bonds. The band at 1734 cm\(^{-1}\) could be due to C=O stretching vibration in the carbonyl groups from terpenoids and flavonoids. The peaks of 1616 cm\(^{-1}\) might indicate presence of C=C bonds or aromatic rings attributed to carbonyl stretch in proteins [41]. The absorption bands appearing at 1244, 1161, 1097 and 1027 cm\(^{-1}\) can be assigned to the C-H alkene (Aliphatic amines, C-H wag, C-N stretching in amide functional group) and C-O vibrations of ether linkages [14, 40, 42]. Thus FTIR analysis of A. indica mediated AgNPs confirmed the presence of different phytochemical components such as flavonoids, polyphenols and terpenoids of A. indica extract which potentially played a vital role in the reduction of Ag\(^{+}\) ions to Ag\(^{0}\) atoms. Especially the proteins (due to the carbonyl and NH\(_2\) of amino acid) present in A. indica leaves extract served as reducing and encapsulating
In order to study the antilarval activity of prepared samples; aqueous extracts of *A. indica* and *C. colocynthis* and AgNPs mediated by these extracts at different concentrations, were subjected separately to larvicidal bioassay on fourth instars larvae of dengue (*A. aegypti*). After 24h treatment, dead and alive mosquito larvae were counted to calculate percent mortality. Larvae which showed no motility after being disturbed with a needle were considered dead. The typical demonstration of larvicidal bioassay are exhibited in figure 11 (*A. indica* panel) and figure 12 (*C. colocynthis* panel).

For *A. indica* leaves aqueous extract, three concentrations; 1 ml, 2 ml and 3 ml of as-prepared extract was tested while five concentrations 1.25 ppm, 2.5 ppm, 5ppm, 10 ppm and 20 ppm of *A. indica* extract mediated AgNPs were exposed to *A. aegypti*. DI water was taken as control in each experiment. In the initial stage of this bioassay at t= 0 h, as shown in upper panel of figure 11, random scattering of the larvae indicate their active movement in all the samples. After 24 h, lower panel of figure 11, all larvae in the control (water) were alive and moved energetically. Beakers containing extract of *A. indica* showed no mortality in all three concentration samples. Nonetheless larvae were found less motile in the sample with 3 ml concentration of extract as compared to other two samples (1ml and 2 ml extract). Beakers containing AgNPs showed different mortalities with different concentrations. The dead larvae can be observed at the bottom or gathered in the middle of the each beaker with high concentration (lower panel of figure 11). AgNPs with concentration of 1.25 ppm, 2.5 ppm, 5ppm, 10 ppm and 20 ppm caused 59%, 66%, 80% and 85% and 100% mortality respectively. All the larvicidal results of *A. indica* leaves extract and mediated AgNPs are summarized in table 1.

**Table 2 Larvicidal activity of DI water (control), aqueous extract of *A. indica* leaves and extract mediated AgNPs at different concentrations against fourth instars larvae of *A. aegypti***.

| Sample type                  | Concentration of sample | Percentage mortality (%) |
|------------------------------|--------------------------|--------------------------|
| Water (control)              | -                        | 0 (active)               |
| as –prepared                 | 1 ml                     | 0 (active)               |
| aqueous extract of *A. indica* leaves | 2 ml                     | 0 (active)               |
| *A. indica* extract mediated AgNPs | 3 ml                     | 0 (less motile)          |
| *A. indica* extract          | 1.25 ppm                 | 49±4                     |
|                              | 2.5 ppm                  | 66±6                     |
|                              | 5 ppm                    | 80±4                     |
|                              | 10 ppm                   | 85±4                     |
|                              | 20 ppm                   | 100±0                    |
A similar larvicidal activity was conducted for samples of *C. colocynthis* fruit extract; in this case again three concentration of as-prepared aqueous extract; 1 ml, 2 ml and 3 ml and DI water as control were taken while for extracted mediated AgNPs five concentration were chosen as 0.3 ppm, 0.6 ppm, 1.25 ppm, 2.5 ppm and 5 ppm (figure 12). In the beginning of the activity test (at 0 h), all larvae were alive and motive in samples (upper panel of figure 12). After exposure of 24 h, again no mortality in (control) and all three extract sample, however, a concentration dependent effect on motility of larvae was noticed in extract samples. The larvae in sample with 3 ml extract were least motile. In the case of AgNPs samples, highest mortality was counted in the sample with maximum concentration. A mortality of 53%, 64%, 80%, 91% and 100% were recorded for samples with AgNPs concentration of 0.3 ppm, 0.6 ppm, 1.25 ppm, 2.5 ppm and 5 ppm respectively. Assembly of immotile (dead) larvae can be witnessed in middle of beakers (lower panel, figure 12). The statistics of *C. colocynthis* mediated larvicidal assay are listed in table 2.

| Sample Type                | Concentration of sample | Percentage Mortality (%) |
|----------------------------|-------------------------|--------------------------|
| Water                      | -                       | 0 (active)               |
| as –prepared aqueous       | 1 ml                    | 0 (active)               |
| extract of *C. colocynthis*| 2 ml                    | 0 (less motile)          |
| fruit                      | 3 ml                    | 0 (least motile)         |
| *C. colocynthis* extract   | 0.3 ppm                 | 53±4                     |
| mediated AgNPs             | 0.6 ppm                 | 64±6                     |
|                            | 1.25 ppm                | 80±3                     |
|                            | 2.5 ppm                 | 91±3                     |
|                            | 5 ppm                   | 100±0                    |

Table 3 Larvicidal results of DI water (control), aqueous extract of *C. colocynthis* fruit and extract mediated AgNPs at different concentrations against fourth instars larvae of *A. aegypti*.

In our case, larvicidal efficacy of both types of aqueous extracts (*A. indica* leaves and *C. colocynthis* fruit) was noticed very insignificant as compared to AgNPs mediated by these extracts. Some other researchers have also reported negligible biocidal activity of aqueous extracts in comparison to extracts mediated nanoparticles [21, 23, 47]. In other hands, a concentration-dependent biocidal efficiency of pure aqueous extracts of various natural products can also be found in literature [16, 28]. It seems that biocidal activities of extracts depend not only on the concentrations used in the bioassays but also on the level of purification (crude, highly purified, aqueous, or alcoholic) as well as nature and different parts of bio-materials to be extracted such as leave, fruit, stem and seed [16, 19, 46].

In both types of extract mediated AgNPs, We observed that by increasing the concentration of AgNPs, percent mortality of larvae increased. Nevertheless, *C. colocynthis* fruit extract mediated AgNPs were found more effective at lower concentrations and exhibited strong larvicidal efficacy as compared to *A. Indica* leaves extract mediated AgNPs as demonstrated graphically in figure 13. As mentioned above, different types of biomaterials (plants) and even various parts of same plant may affect the biocidal efficiency of mediated AgNPs. That’s why perhaps AgNPs prepared by fruit extract of *C. colocynthis* showed better antilarval activity even at lower concentrations in our case. A comparative larvicidal study of extracts and mediated AgNPs using various parts such as leaves, fruit, seed and stem of same plant can be conducted for more detailed investigations.

Since the exact mechanism of larvae mortality is unidentified, different modes of action can be proposed for larvicidal activity of AgNPs. Most probably the size and surface chemistry of AgNPs play a vital role to make them lethal against mosquito larvae. At the first stage, owing to small size AgNPs penetrate into the larval membrane causing the possible leakage of cellular materials. In the next step, AgNPs may interact with the different cell molecules to inhibit molting and other physiological processes. In their action, AgNPs may produce peroxide, inactivate the enzymes and disturb the functional proteins leading to the larvae death. The presence of extract biomolecules on the AgNPs surface can further enhance their effectiveness in this larvicidal process [23, 40, 46, 47]. This indicates that anti-dengue pharmaceuticals can be formulated using very low concentration of natural products based AgNPs for practical applications. Thus findings of this study can be helpful to provide new paradigm in designing the green chemistry based alternative to combat the increasing mosquitos’ diseases.
5. Conclusion

AgNPs were prepared from herbal origin which is economical and eco-friendly using the aqueous extracts of *A. indica* leaves and *C. colocynthis* fruit. Both types of NPs were found pure metallic silver in nature having FCC crystalline structure and of spherical in shapes with diameters in the range of 25±5 nm, synthesized by different biomolecules of extracts as determined by different characterization techniques including SEM, UV-Vis, XRD and FTIR. The prepared green AgNPs exhibited great larvicidal potential against *A. aegypti* even at low concentrations; LC$_{50}$ at 1.25 ppm for *A. indica* and LC$_{50}$ at 0.3 ppm *C. colocynthis* mediated AgNPs respectively. We observed a concentration dependent larvicidal activity in both types of AgNPs whereas their extracts showed negligible efficiency in both cases. These results indicate the potential use of green AgNPs as alternative to replace the synthetic products in pharmaceuticals for anti-dengue purposes.

Acknowledgments

Authors acknowledge Dr. Shahid Atiq and Dr. Syed Sajjad Hussain from Centre of Excellence in Solid State Physics, University of the Punjab, Lahore- Pakistan for their cooperation in experimental work and useful discussions in manuscript writing.

Ethical Statement

It is not relevant to our work.

Funding Statement

This work was financially supported by Higher Education Commission (HEC) of Pakistan under the Project of ‘National Research Program for Universities’ (Project No.: HEC-NRPU-8019).

Data Accessibility

Our data are deposited at Dryad: [https://datadryad.org/stash/dataset/doi:10.5061/dryad.z612jm68k](https://datadryad.org/stash/dataset/doi:10.5061/dryad.z612jm68k), with following reviewer URL:

Dryad Reviewer URL: [https://datadryad.org/stash/share/DksW82FCdqlSwPql2aYhDFl4C9QUE74hEBPTmrJ3trHw](https://datadryad.org/stash/share/DksW82FCdqlSwPql2aYhDFl4C9QUE74hEBPTmrJ3trHw)

Competing Interests

We have no competing interests.

Authors’ Contributions

1). SR and MAR performed the synthesis experiments, analysed the data and wrote the manuscript
2). ZK, SR and MAR designed the experiments and conducted the larvicidal bioassays
3). SR and SN conducted the characterization measurements and helped in interpretation of data
4). FM and ZK provided the larvae facility and helped in larvicidal activity experiments
5). SR and SN conducted the characterization measurements and helped in interpretation of data
6). MJR and SR conducted the FTIR and SEM characterization and helped in data analysis

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Figure 1. Different steps during the preparation of aqueous extract of *A. indica* leaves; (a) fresh leaves were collected, (b) leaves were kept under shade and dried on aluminum sheet (c) boiling of very small pieces *A. indica* leaves in DI water (d) *A. indica* extract in sample bottle after filtration

Figure 2. Different stages of *C. colocynthis* fruit extract preparation; (a) green fruit was cut into very small pieces, (b) small pieces soaked into DI water for 72 h at room temperature, (c) finally aqueous extracts were obtained by filtration.
Figure 3. (Upper panel) different phases of the synthesis of A. indica extract mediated AgNPs. (a) Formation of Ag-ions in the colorless precursor AgNO₃ solution during heating, (b) dropwise adding of extract during heating and constant stirring appearing of color indicate the reduction of Ag-ions, (c) Final stage color indicating the complete reduction of Ag-ions into AgNPs by the extract. (Lower panel) different samples in the bottles (a) transparent AgNO₃ solution, (b) yellowish green aqueous extract of A. indic, (c) colloidal sample of AgNPs mediated by 1000µl of extract, (d) AgNPs prepared by 500µl extract, and (e) AgNPs synthesized by 250µl of extract.

Figure 4. (upper panel) Different levels during the formation of AgNPs using the aqueous extract of C. colocynthis (a) transparent precursor solution, (b) solution after mixing the extract by stirring and heating, a slight color change to light yellowish brown, (c) further evident color change with time during heating and stirring indicate the reduction of Ag-ions to Ag-atoms and formation of AgNPs (d) completion of the reaction with reddish brown color of AgNPs. (lower panel) (a) AgNO₃ colorless precursor solution, (b) light yellow color C. colocynthis aqueous extract, (c-d) colloidal sample of AgNPs prepared by 4ml, 2.5 ml and 1 ml of C. colocynthis extract respectively.

Figure 5. Schematic illustration of synthesis mechanism of extract mediated AgNPs. The phenomena occurring in each phase in mentioned accordingly.

Figure 6. UV-Vis spectroscopy results of AgNPs synthesized by using; (a) three different concentrations of A. indica leaves extract, (b) three different concentration of C. colocynthis fruit aqueous extract. Single absorption peak in each case appearing in the wavelength range of 412 nm to 430 nm indicate the spherical shaped nano-sized silver particles. Insets display images of each colloidal sample in cuvette.

Figure 7. SEM micrographs and size distribution histograms of both types of biosynthesized AgNPs; (a) SEM image of A. indica mediated AgNPs showing the almost spherical shaped NPs decorated on the larger sized extract particles, (b) particle size histogram of A. indica AgNPs indicating the diameter range 12-24 nm. (c) SEM results of C. colocynthis mediated AgNPs revealing the sphere like morphology and nice random dispersal over the bigger extract particles. The average size of these AgNPs was found to be 26± 5 nm as can be seen in the histograms (d).

Figure 8. EDX spectrum showing elemental composition analysis of AgNPs prepared by the aqueous extracts of (a) A. indica leaves and (b) C. colocynthis fruit. Insets show the table of elements in each case. The presence of elemental silver and other biomaterials can be discerned by different absorption characteristics peaks in the both spectra.

Figure 9. X-ray diffraction patterns of (a) A. indica mediated AgNPs and (b) C. colocynthis mediated AgNPs. In both cases, along with characteristics peaks designating the face centered cubic (fcc) crystalline nature of prepared nanoparticles, appearance of few extra peaks (mentioned by stars *) indicate the presence of some extract molecules.

Figure 10. FTIR spectra of both types of AgNPs; (a) A. indica leaves extract mediated AgNPs, (b) AgNPs prepared using extract of C. colocynthis fruit. Occurrence of various absorption peaks at different positions in each case indicates the presence of difference biomolecules on the surface of synthesised nanoparticles.

Figure 11. A representative display of larvicidal bioassay for different test samples in the beakers including water control (water), A. indica extract and its mediated AgNPs at different concentrations (as mentioned); upper panel shows the initial stage of experiment with 0h time while results after 24 h are represented in lower panel. It can be noticed that in the beginning all larvae were active and randomly scattered but after 24h due to mortality most of the dead larvae are gathered in the middle of the beakers. However, in control, extract and samples with low AgNPs concentration scattered larvae can be discerned.

Figure 12. A typical demonstration of larvicidal activity of C. colocynthis extract mediated AgNPs with different concentrations, as prepared extract and water as control. In upper panel, different samples with larvae are shown in the start of experiment (at 0h) where random sprinkle of the larvae into the liquid can be seen. After 24h of exposure, larvicidal efficacy of different samples is presented in the lower panel where dead larvae can be found assembled at the bottom of the beakers.
Figure 13. Graphs showing the larvicidal performance in mortality (%) of both types of AgNPs at different test concentrations; (A) *A. indica* extract mediated AgNPs showed LC$_{50}$ at about 1.25 ppm while at 20 ppm 100% mortality was observed, (B) AgNPs prepared by *C. colocynthis* extract exhibited 50% mortality (LC$_{50}$) only at about 0.3 ppm and 100% mortality at 5ppm.

Figures

![Figure 1](image1.png)

![Figure 2](image2.png)
Figure 3

Figure 4
Figure 5

1. Silver nitrate (AgNO₃) creation of silver ions (Ag⁺)
2. Reduction process Ag⁺ → Ag⁰
3. Formation of silver nuclei
4. Nucleation and Growth processes to develop AgNPs

Figure 6

(a) A.indica mediated AgNPs
(b) C.colocynthis mediated AgNPs
Figure 7

![Figure 7](image)

**Elemental Analysis**

| Element | Standard Label | Weight (%) |
|---------|----------------|------------|
| C       | C              | 38.94      |
| O       | SiO₂           | 15.93      |
| Na      | NaK₂O₂         | 12.12      |
| Mg      | MgO            | 0.58       |
| Si       | SiO₂           | 4.87       |
| Cl       | Cl             | 1.14       |
| K       | K              | 1.14       |
| Ag       | Ag             | 1.14       |
| Au       | Au             | 1.14       |

Total: 100

Figure 8

![Figure 8](image)

**Elemental Analysis**

| Element | Standard Label | Weight (%) |
|---------|----------------|------------|
| C       | C              | 31.87      |
| O       | SiO₂           | 15.93      |
| Na      | NaK₂O₂         | 12.12      |
| Mg      | MgO            | 0.58       |
| Si       | SiO₂           | 4.87       |
| Ca      | CaSiO₃         | 1.14       |
| K       | K              | 1.14       |
| Ag       | Ag             | 1.14       |
| Au       | Au             | 1.14       |

Total: 100
Figure 12

Figure 13
Figure 1. Different steps during the preparation of aqueous extract of A. indica leaves; (a) fresh leaves were collected, (b) leaves were kept under shade and dried on aluminum sheet (c) boiling of very small pieces A. indica leaves in DI water (d) A. indica extract in sample bottle after filtration.

227x71mm (150 x 150 DPI)
Figure 2. Different stages of C. colocynthis fruit extract preparation; (a) green fruit was cut into very small pieces, (b) small pieces soaked into DI water for 72 h at room temperature, (c) finally aqueous extracts were obtained by filtration.

194x77mm (150 x 150 DPI)
Figure 3. (Upper panel) different phases of the synthesis of A. indica extract mediated AgNPs. (a) Formation of Ag-ions in the colorless precursor AgNO₃ solution during heating, (b) dropwise adding of extract during heating and constant stirring appearing of color indicate the reduction of Ag-ions, (c) Final stage color indicating the complete reduction of Ag-ions into AgNPs by the extract. (Lower panel) different samples in the bottles (a) transparent AgNO₃ solution, (b) yellowish green aqueous extract of A. indic, (c) colloidal sample of AgNPs mediated by 1000µl of extract, (d) AgNPs prepared by 500µl extract, and (e) AgNPs synthesized by 250µl of extract.

215x227mm (150 x 150 DPI)
Figure 4. (upper panel) Different levels during the formation of AgNPs using the aqueous extract of *C. colocynthis* (a) transparent precursor solution, (b) solution after mixing the extract by stirring and heating, a slight color change to light yellowish brown, (c) further evident color change with time during heating and stirring indicate the reduction of Ag-ions to Ag-atoms and formation of AgNPs (d) completion of the reaction with reddish brown color of AgNPs. (lower panel) (a) AgNO₃ colorless precursor solution, (b) light yellow color *C. colocynthis* aqueous extract, (c-d) colloidal sample of AgNPs prepared by 4ml, 2.5 ml and 1 ml of *C. colocynthis* extract respectively.

254x193mm (150 x 150 DPI)
Figure 5. Schematic illustration of synthesis mechanism of extract mediated AgNPs. The phenomena occurring in each phase is mentioned accordingly.

286x153mm (150 x 150 DPI)
Figure 6. UV-Vis spectroscopy results of AgNPs synthesized by using; (a) three different concentrations of *A. indica* leaves extract, (b) three different concentration of *C. colocynthis* fruit aqueous extract. Single absorption peak in each case appearing in the wavelength range of 412 nm to 430 nm indicate the spherical shaped nano-sized silver particles. Insets display images of each colloidal sample in cuvette.
Figure 7. SEM micrographs and size distribution histograms of both types of biosynthesized AgNPs; (a) SEM image of A. indica mediated AgNPs showing the almost spherical shaped NPs decorated on the larger sized extract particles, (b) particle size histogram of A. indica AgNPs indicating the diameter range 12-24 nm. (c) SEM results of C. colocynthis mediated AgNPs revealing the sphere like morphology and nice random dispersal over the bigger extract particles. The average size of these AgNPs was found to be 26± 5 nm as can be seen in the histograms (d).

237x187mm (150 x 150 DPI)
Figure 8. EDX spectrum showing elemental composition analysis of AgNPs prepared by the aqueous extracts of (a) A. indica leaves and (b) C. colocynthis fruit. Insets show the table of elements in each case. The presence of elemental silver and other biomaterials can be discerned by different absorption characteristics peaks in both spectra.

| Elements | Standard label | Weight (%) |
|----------|----------------|------------|
| C        | C Vitr        | 38.99      |
| O        | SiO₂          | 11.38      |
| Na       | Na₅Si₅O₁₂     | 0.26       |
| Mg       | MgO           | 0.17       |
| Si       | SiO₂          | 1.42       |
| Cl       | NaCl          | 4.62       |
| K        | KBr           | 2.3        |
| Ag       | Ag            | 37.08      |
| Au       | Au            | 3.77       |
| Total    |               | 100        |

| Element  | Standard Label | Weight (%) |
|----------|----------------|------------|
| C        | C Vitr        | 31.87      |
| O        | SiO₂          | 15.93      |
| Na       | Na₅Si₅O₁₂     | 0.65       |
| Mg       | MgO           | 0.36       |
| Si       | SiO₂          | 4.87       |
| Ca       | CaSi₅O₁₂      | 1.1        |
| Cl       | NaCl          | 1.16       |
| K        | KBr           | 3.1        |
| Pd       | Pd            | 0.58       |
| Ag       | Ag            | 38.5       |
| Au       | Au            | 3.87       |
| Total    |               | 100        |
Figure 9. X-ray diffraction patterns of (a) A. indica mediated AgNPs and (b) C. colocynthis mediated AgNPs. In both cases, along with characteristics peaks designating the face centered cubic (fcc) crystalline nature of prepared nanoparticles, appearance of few extra peaks (mentioned by stars *) indicate the presence of some extract molecules.
Figure 10. FTIR spectra of both types of AgNps; (a) A. indica leaves extract mediated AgNPs, (b) AgNPs prepared using extract of C. colocynthis fruit. Occurrence of various absorption peaks at different positions in each case indicates the presence of difference biomolecules on the surface of synthesised nanoparticles.
Figure 11. A representative display of larvicidal bioassay for different test samples in the beakers including water control (water), A. indica extract and its mediated AgNPs at different concentrations (as mentioned); upper panel shows the initial stage of experiment with 0h time while results after 24 h are represented in lower panel. It can be noticed that in the beginning all larvae were active and randomly scattered but after 24h due to mortality most of the dead larvae are gathered in the middle of the beakers. However, in control, extract and samples with low AgNPs concentration scattered larvae can be discerned.
A typical demonstration of larvicidal activity of *C. colocynthis* extract mediated AgNPs with different concentrations, as prepared extract and water as control. In upper panel, different samples with larvae are shown in the start of experiment (at 0h) where random sprinkle of the larvae into the liquid can be seen. After 24h of exposure, larvicidal efficacy of different samples is presented in the lower panel where dead larvae can be found assembled at the bottom of the beakers.
Figure 13. Graphs showing the larvicidal performance in mortality (%) of both types of AgNPs at different test concentrations; (A) A. indica extract mediated AgNPs showed LC50 at about 1.25 ppm while at 20 ppm 100% mortality was observed, (B) AgNPs prepared by C.colocynthis extract exhibited 50% mortality (LC50) only at about 0.3 ppm and 100% mortality at 5ppm.
Appendix B

Reply to Reviewer’s Reports

(Manuscript ID RSOS-200540)

We are thankful to the reviewers for their expert assessment of our manuscript. We are grateful for their thoughtful and valuable comments to improve our manuscript. All points and issues raised by the learnt reviewers have been considered and the manuscript has been modified accordingly. A detailed point-by-point response to all comments is provided below indicating the implemented changes in the revised version of the manuscript. A highlighted version by ‘Track Changes’ is also included with the resubmission.

Response to 1st Reviewer’s Comments

Comment 1. The article describes the synthesis of Ag nanoparticles using *Azadirachta indica* and *Citrullus colocynthis* extracts. The work done is useful and is in line with the current scientific demands. There are also some issues to be addressed before publishing this work.

Response 1. We are grateful to the learnt reviewers for mentioning our work “useful “and “in line with current scientific demands”.

Comment 2. There are several grammatical mistakes for instance I am highlighting some of these:

i) page 2 line 17 which have famous is incorrect it should be which are famous

ii) page 2 line 25 “vector for many deceases like” it should be vector for many diseases like

iii)page 2 line 25 please correct “fever.Dengue”

iv) page 2 line 35 “AgNPs were manufactured” were synthesized

v) page 2 line 35 “the mixture was let to cool down to room temperature” should be the mixture was cooled to room temperature

The authors should thoroughly check the whole manuscript and English should be improved as well as all the editorial mistakes should be rectified.
Response 2. We are thankful to the reviewer for pointing out these grammatical mistakes to improve beauty of the text. The whole manuscript has been thoroughly checked and all the grammatical and editorial mistakes mentioned by the reviewers and otherwise are corrected.

Comment 3. Page 6 line 10, authors mentioned the particle size, I guess it should be crystallite size?

Response 3. The corrections were made as suggested by the reviewer.

Comment 4. Authors have not mapped the particle size in SEM images. The particle size should be measured in SEM image and it should be visible in the picture.

Response 4. According to the reviewer recommendations, the SEM images have been modified by adding the new images showing particle size visible in the picture.

Comment 5. The authors have mentioned that the chosen plants have known biological activity. However, they do not show any activity against A. aegypti. Secondly, if the extracts are not active against A. aegypti why there is a large difference in activity of AgNPs prepared from two different extracts as LC50 at 1.25 ppm for A. indica and LC50 at 0.3 ppm for C. colocynthis is observed. The later is 4 times more active than the former one. I do not seem that activity is only due to AgNPs in that case it should be similar or close to each other.

Response 5. We are appreciative to the reviewer for highlighting this point. Firstly, it is well-established fact that various medicin plants including A. indica and C. colocynthis, exhibit different biological activities (R1-R3). However, in our case, both of the aquous extracts (A. indica leaves and C. colocynthis fruit) did not show any larvicidal efficacy against dengue vector. The inefficiency of aquous extracts of different plants was also reported by other researchers [R3-R6] however, some studies report concentration-dependent biological efficency of aquous extracts of some natural products [R2,R7]. Thus, biological activity of extracts of biomaterias depend on various
factors such as nature, different parts (leave, fruit, stem and seed) and level of purification (crude, highly purified, aqueous, or alcoholic) and concentration [R8-R9].

The reason of the difference in larvicidal activity of AgNPs prepared by A. indica leaves or C. colocynthis fruit is that different types of biomolecules are deposited on the surface of AgNPs (as confirmed by FTIR analysis). This is, in fact, the synergistic effect of Ag-ions and biomolecules to enhance the larvicidal performance of extract mediated AgNPs. Thus the difference in larvicidal efficiency occurs due to the presence of different types of biomolecules on the surfaces of AgNPs although Ag-ions are same in both cases. Therefore, the LC$_{50}$ values for both cases were different (LC$_{50}$ at 1.25 ppm for A. indica and LC$_{50}$ at 0.3 ppm for C. colocynthis).

Main text has also been modified to explain this point.

**Comment 6.** Figures 1, 2 and 5 are unnecessary and should be deleted.

**Response 6.** Figures 1 and 3 and figures 2 and 4 are combined according to the recommendations of reviewer 2, instead of deleting.

Figure 5 depicts schematics of the potential formation mechanism of extract mediated AgNPs and different phases of nucleation and growth processes. We would like to keep it in the main text with the kind consent of our proficient reviewer.

Data of all modified figures has been uploaded to Dryad Digital Repository

Our data are deposited at Dryad:

https://datadryad.org/stash/dataset/doi:10.5061/dryad.z612jm68k

Dryad Reviewer URL:

https://datadryad.org/stash/share/DksW82FCDqISwPqI2aYhDl4C9QUE74hEBPTmrJ3trHw

**Comment 7.** the sentence in conclusion section “Both types of NPs were found pure metallic silver in nature having FCC crystalline structure and of spherical in shapes with diameters in the range of 25±5 nm, synthesized by different biomolecules of extracts as determined by different characterization techniques including SEM, UV-Vis, XRD and FTIR” is too long and understandable. It should be refabricated to two sentences.

After addressing these issues the article can be published
Response 7. We are agreed with the reviewer and this sentence has been rephrased into two sentences as,
“Both types of NPs were found pure metallic silver in nature having FCC crystalline structure and were spherical in shapes with diameters in the range of 25±5 nm, as determined by SEM and XRD. The optical behavior of colloidal samples and presence of different extract biomolecules were studied by UV-Vis, and FTIR.” The main text has also been modified accordingly

We are obliged to the reviewer for recomemnding our manuscript for “publication” in RSOS.

Response to 2nd Reviewer’s Comments

Comment 1. The MS presents work which has been published with various other plants. The work carried out is significant but need major revision.

Response 1. We are obliged to the reviewer for mentioning our work “significant”.

Comment 2. MS needs enormous revision in scientific reporting, language and grammar. The scientific names are wrongly spelled at some places.

Response 2. We are grateful to the learnt reviewer for highlighting the errors to enhance the flouncy and beauty of the text. The manuscript is carefully examined and all mistakes of language, grammar, typos and scientific reporting including scientific names pointed out by the reviewer and otherwise are corrected and highlighted by ‘Track Changes’ in the revised manuscript.

Comment 3. Each section of the MS needs major revision. The comments have been marked on the MS. Authors are advised to go through each and every comment; and rectify accordingly.

Response 3. We are thankful to the reviewer for valuable comments and suggestions. We have addressed all the comments and points raised by the learnt reviewer and
manuscript has been thoroughly revised in the light of remarks and recommendations of the reviewer. The detailed response to the comments is described below to highlight the amendments made in the revised version of the manuscript accordingly.

**Comment:** The MS is about anti-dengue vector. Kindly revise.

**Response:** The title of the MS has been rectified as directed by the reviewer.

**Comment:** Check series.... FTIR was done in last..Please write based on conducting the characterization

**Response:** In abstract, series of the characterizations techniques has been corrected, as suggested by the reviewer.

**Comment:** Authors have not added any comparison between the characteristics of two kinds of NPs. Some features may be mentioned.

**Response:** We agreed to the reviewer and the abstract has been modified by adding comparative characteristic features of both types of AgNPs.

**Comment:** write scientific names correctly throughout the manuscript. Start specific names with small case. Take care of spellings. A. indica and C. colocynthis

**Response:** All scientific names and terms have been corrected including A. indica and C. colocynthis, as directed by the reviewer.

**Comment:** Which part? As you have prepared NPs with fruits, please specify

**Response:** The C. colocynthis fruit has been specified and reference has also been added in the main text, as suggested by the reviewer.

**Comment:** Dengue is transmitted by Aedes mosquitoes. It is caused by virus. Larvae have no role in it. Please rectify Write as hyphenated throughout the MS
Response: We agreed to the reviewer and the sentence has been rectified in the main text as; “Dengue fever incidence has increased 30-fold worldwide in the last 30 years which is caused by dengue virus.”

Comment: What do you mean by glass substrate?

Response: Glass substrate means just a small piece of glass slide on which few drops of colloidal AgNPs sample are dried to make a thick enough NPs layer. This was used as sample for characterization purposes. It has also been described in literature [R10, R11].

Comment: 1ml, 2.5 ml and 4ml.................. on what basis the ratios (pattern of conc.) were decided? The amount/conc. of a compound to be used in a mixture is always in a particular ratio. For example...1, 2, 4, 8....

However, only AgNPs prepared by using the 500 µl of extract was used in larvicidal activity and other characterization, specify the reason plz

Response: In this study, two types of extracts were used as reducing and stabilizing agents for synthesis of AgNP. The amount of extracts in both cases was selected from the literature protocols [R12, R13]. To obtain the optimized properties three different amounts of each extract were used for the preparation of AgNPs. An amount was selected and then its 2 fold and 4 fold was used for AgNPs synthesis process. In the case of A. indica extract, 250 µL, 500 µL and 1000µL while for C. colocynthis extract 1 mL, 2 mL and 4 mL were used (2.5 mL was typo error which has been corrected in the main text). The AgNPs prepared from 500 µL (A. indica) and 2 mL (C. colocynthis) were taken as optimized AgNPs on the basis of Uv-Vis results. This point has been addressed in the main text according to the reviewer’s recommendations.

Comment: Why did authors use directly 5 mM concentration to prepare NPs? Higher concentration of silver nitrate itself is toxic? Did you try for lower concentrations? What was the result? If not, then why?
Response: We agreed to the reviewer that AgNPs can be made by different concentrations of precursor AgNO₃ solution (Lower and higher than 5mM). In this study, both types of AgNPs were prepared following the protocols mentioned in the literature. *A. indica* extract-mediated AgNPs were synthesized using 1mM solution of AgNO₃ using methods described by of Aparajita et al. [R12] while *C. colocynthis* extract-mediated AgNPs using 5mM concentration of precursor solution as reported by Shawkey et al. [R13].

Comment: We cannot say that the fruit extract was more effective at lower concentration as the comparison is between 1 mM and 5 mM of AgNO₃. 5 mM is itself a very high concentration of silver nitrate. If authors want to compare the efficacy then they should do at similar concentrations either 1 mM or 5 mM.

Response: In our work we used two different types of extracts and AgNPs to study the larvicidal activity. Each type of AgNPs were prepared following different protocols using different precursor concentration (1mM or 5mM) and two different reducing agents (*A. indica* leaves extract or *C. colocynthis* fruit extract). We aim to report the antilarvae performance of each type of extract and AgNPs at different concentrations. It is well-known fact that change in concentration of precursor solution will affect the size and morphology of prepared AgNPs [R14, R15]. In our case, we used two type of extracts as reducing and stabilizing agents and thus different types of biomolecules were deposited on the surface of each type of AgNPs which caused variation in larvicidal efficacy.

We have rectified the main text (abstract, result and discussion and conclusions) to address the comparison issue raised by the reviewers.

Comment: ppm unit is no longer in use. May be replaced

Response: According to reviewer recommendations, ppm has been replaced by mg/L throughout the manuscript.

Comment: “The test concentrations of both types of AgNPs were selected on the basis of pilot experiments in which LC50
concentrations were found”. Any published result? If so, give ref. Otherwise, you may give values obtained here.

**Response:**

The LC$_{50}$ values of pilot experiments were found to be 1.25 mg/L and 0.3 mg/L for *A. indica* and *C. colocynthis* mediated AgNPs respectively. These LC$_{50}$ values have also been mentioned in the main text according to the reviewer suggestions.

**Comment:**

This time dependent color change during the synthesis, actually, ........ time not mentioned.

**Response:**

Time has been mentioned in the main text as “10 min for *A. indica* mediated AgNPs and 20 min for *C. colocynthis* mediated AgNPs.”

**Comment:**

we can notice strong silver signal along with some other signals. mention Ag%

**Response:**

% age values of Ag have been mentioned for both types of AgNPs in main text as “37.08 % for *A. indica* AgNPs and 38.5% for *C. colocynthis* AgNPs” as directed by the reviewer.

**Comment:**

mL denotes the volume. As per WHO protocol, 1 mL of toxicant is added to 249 mL of water to study larval mortality. What about concentration? Write the conc of extract. How can you compare the mL of extract with ppm of AgNPs?

**Response:**

The reviewer is right that ml determines the volume only. We have added percentage concentration of the extract. It is in common practice to use the extract in percentage concentration [R16, R17]. The prepared extract is standardized as 100 % from which dilutions are drawn for biassays. We used 3 different amounts (1mL, 2mL and 3mL) of the extract to be dissolved in water to make a final volume of 200 mL. 1 mL of extract in 200 mL of water will make a 0.5 % extract concentration, 2 mL will make 1 % and 3 mL will make 1.5 % extract concentration in the solution. Furthermore because the dynamics of the concentration formulations for extract and nanoparticles is different, extract is based on the extracted material while nanoparticles’ concentration is
devised from the precursor used, therefore, it is not necessary to have a direct comparison of the two in terms of concentration units.

**Comment:** 1ml, 2ml, 4 ml............ Are these concentrations? , 2, 3 mL..... Is this concentration?

**Response:** No, these are quantities, now we have added % age concentration of extracts (0.5%, 1%, and 1.5%) in main text to rectify this issue raised by the reviewer.

**Comment:** How can you make anti-dengue medicine which has been tested against dengue larvae not dengue virus

**Response:** We are thankful to the reviewer for indicating this point, we have rectified the main main text as “This indicates that antilarval pharmaceuticals can be formulated using very low concentration of natural products based AgNPs for practical applications”

**Comment:** Figure captions: all figure captions need to be revised...as they too lengthy, unnecessarily too big legends. Please make all legends concise and crisp.

**Response:** All the figure captions have been revised to make brief and concise, as directed by reviewer.

**Comment:** Figs. 1, 2 and 3 can be easily combined. Please shorten and make it crisper. F1: explanations can be included in the figures itself (labellings of figures)

**Response:** According to the reviewer recommendations, previous figure 1 and 3 have been combined into one (new figure no. 1) and figure 2 has been combined with figure 4 to make new figure no. 2. The main text has also been amended accordingly.

**Comment:** F11 and F12: Please delete...instead add LC values tables
Response: The LC values have already been added in tables 1 and 2. The figures 11 and 12 provide the visual demonstration of the larvicidal activity, so with the kind consent of our proficient reviewer we would like to keep them in the main text.

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R3. Govindachari TR, Suresh G, Gopalakrishnan G, Banumathy B, Masilamani S. 1998 Identification of antifungal compounds from the seed oil of Azadirachta indica. Phytoparasitica 26, 109–116. (doi: 10.1007/BF02980677)

R4. Shawkey AM, Abdulall AK, Rabeh MA, Abdellatif AO. 2014 Enhanced biocidal activities of Citrullus colocynthis aqueous extracts by green nanotechnology. International Journal of Applied Research in Natural Products 7 (2), 1-10.

R5. Shawkey AM, Rabeh MA, Abdulall AK, Abdellatif AO. 2013 Green nanotechnology: Anticancer activity of silver nanoparticles using Citrullus colocynthis aqueous extracts. Adv. Life Sci. Technol. 13, 60–70.

R6. Rawani A, Ghosh A, Chandra G. 2013 Mosquito larvicidal and antimicrobial activity of synthesized nano-crystalline silver particles using leaves and green berry extract of Solanum nigrum L. (Solanaceae: Solanales). Acta Tropica 128, 613–622. (doi: 10.1016/j.actatropica.2013.09.00)

R7. Sujitha V, Murugan K, Paulpandi M, Panneerselvam C, Suresh U, Roni M, Nicoletti M, Higuchi A, Madhiyazhagan P, Subramaniam J, Dinesh D, Vadivalagan C, Chandramohan B, Alarfaj AA, Munusamy MA, Barnard DR, Benelli G. 2015 Green-synthesized silver nanoparticles as a novel control tool against dengue virus (DEN-2) and its primary vector Aedes aegypti. Parasitol Res., 114, 3315-25. (doi: 10.1007/s00436-015-4556-2)

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R10. Kanwal Z, Raza MA, Riaz S, Manzoor S, Tayyeb A, Sajid I, Naseem S. 2019 Synthesis and characterization of silver nanoparticle-decorated cobalt nanocomposites (Co@AgNPs) and their density-dependent antibacterial activity. *R. Soc. open sci.* **6**, 182135. (doi:10.1098/rsos.182135)

R11. Kanwal Z, Raza, MA, Manzoor F, Riaz S, Jabeen G, Fatima S, Naseem S. 2019 A Comparative Assessment of Nanotoxicity Induced by Metal (Silver, Nickel) and Metal Oxide (Cobalt, Chromium) Nanoparticles in *Labeo rohita*. *Nanomaterials*, **9**, 309. (doi:10.3390/nano9020309)

R12. Aparajita V, Mohan SM. 2016 Controllable synthesis of silver nanoparticles using Neem leaves and their antimicrobial activity. *J Radiat Res Appl Sc* **9**(1), 109-115. (doi: 10.1016/j.jrras.2015.11.001)

R13. Shawkey AM, Rabeh MA, Abdulall AK, Abdellatif AO. 2013 Green nanotechnology: Anticancer activity of silver nanoparticles using Citrullus colocynthis aqueous extracts. *Adv. Life Sci. Technol.* **13**, 60–70.

R14. Devadiga A, Vidya SK, Saidutta MB. 2017 Effect of Precursor Salt Solution Concentration on the Size of Silver Nanoparticles Synthesized Using Aqueous Leaf Extracts of *T. catappa* and *T. grandis* Linn f.—A Green Synthesis Route. In: Mohan B. R., Srinikethan G., Meikap B. (eds) Materials, Energy and Environment Engineering. Springer, Singapore. (doi: 10.1007/978-981-10-2675-1_17)

R15. Vishwakarma K, Shweta, Upadhyay N, Singh J, Liu S, Singh VP, Prasad SM, Chauhan DK, Tripathi DK, Sharma S. 2017 Differential Phytotoxic Impact of Plant Mediated Silver Nanoparticles (AgNPs) and Silver Nitrate (AgNO₃) on *Brassica sp*. *Front Plant Sci* **8**, 1501. (doi: 10.3389/fpls.2017.01501)

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R17. Ederley V, Gloria C, Gladis M, César H, Jaime O, Oscar A. 2018 Silver Nanoparticles Obtained by Aqueous or Ethanolic *Aloe vera* Extracts: An Assessment of the Antibacterial Activity and Mercury Removal Capability. *J Nanomater* 2018, Article ID 7215210, 7 pages (doi: 10.1155/2018/7215210)
Biosynthesis, Characterization and Anti-dengue vector activity of Silver Nanoparticles prepared by Azadirachta indica and Citrullus colocynthis

| Journal: | Royal Society Open Science |
| --- | --- |
| Manuscript ID | RSOS-200540.R1 |
| Article Type: | Research |
| Date Submitted by the Author: | 04-Jul-2020 |
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| Subject: | Green chemistry < CHEMISTRY, Nanotechnology < CHEMISTRY, biophysics < CROSS-DISCIPLINARY SCIENCES |
| Keywords: | silver nanoparticles, green synthesis, Citrullus colocynthis, Azadirachta indica, antilarval activity |
| Subject Category: | Chemistry |
Author-supplied statements

Relevant information will appear here if provided.

Ethics

Does your article include research that required ethical approval or permits?:
This article does not present research with ethical considerations

Statement (if applicable):
CUST_IF_YES_ETHICS :No data available.

Data

It is a condition of publication that data, code and materials supporting your paper are made publicly available. Does your paper present new data?:
Yes

Statement (if applicable):
Data Accessibility statement:

Our data are deposited at Dryad:
https://datadryad.org/stash/dataset/doi:10.5061/dryad.z612jm68k [48], with following reviewer URL;

Dryad Reviewer URL:
https://datadryad.org/stash/share/DksW82FCDqlSwPql2aYhD14C9QUE74hEBPTmrJ3trHw

Conflict of interest

I/We declare we have no competing interests

Statement (if applicable):
CUST_STATE_CONFLICT :No data available.

Authors’ contributions

This paper has multiple authors and our individual contributions were as below

Statement (if applicable):
Authors’ Contributions:

1). SR and MAR performed the synthesis experiments, analysed the data and wrote the manuscript
2). ZK, SR and MAR designed the experiments and conducted the larvicidal bioassays
3). SR and SN conducted the characterization measurements and helped in interpretation of data
4). FM and ZK provided the larvae facility and helped in larvicidal activity experiments
5). SR and SN conducted the characterization measurements and helped in interpretation of data
6). MJR and SR conducted the FTIR and SEM characterization and helped in data analysis
Biosynthesis, Characterization and Anti-dengue vector activity of Silver Nanoparticles prepared by *Azadirachta indica* and *Citrullus colocynthis*

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Keywords: antilarval-dengue activity, silver nanoparticles, green synthesis, *Azadirachta indica*, *Citrullus colocynthis*

Abstract

We report here biosynthesis of silver nanoparticles (AgNPs) using aqueous extracts of (i) *Azadirachta indica* leaves and (ii) *Citrullus colocynthis* fruit and their larvicidal activity against *Aedes aegypti*. The UV-Vis spectroscopy absorption peaks occurred in the range of 412-416 nm for *A. indica* AgNPs and 416-431 nm for *C. colocynthis* AgNPs indicating the silver nature of prepared colloidal samples. The Scanning electron microscopy examination revealed the spherical morphology of both types of NPs with average size of 17±4 nm (*A. indica* AgNPs) and 26±5 nm (*C. colocynthis* AgNPs). The x-ray diffraction pattern confirmed the face centred cubic (FCC) structure with crystallite size of 11±1 nm (*A. indica* AgNPs) and 15±1 nm (*C. colocynthis* AgNPs) while characteristic peaks appearing in Fourier transform infrared spectroscopy analysis indicated the attachment of different biomolecules on AgNPs. The larvicidal activity at different concentrations of synthesized AgNPs (1-20 mg/L) and extracts (0.5-1.5 %) against *Aedes aegypti* was examined for 24 h. A concentration dependent larvicidal potential of both types of AgNPs was observed. The LC₅₀ values were found to be 0.3 mg/L and 1.25 mg/L for *C. colocynthis* AgNPs and *A. indica* AgNPs respectively. However, both extracts did not exhibit any notable larvicidal activity.

1. Introduction

Nanotechnology, owing to unique, extraordinary and incredible properties of materials at nano-scale, has revolutionized almost every field of science and technology. Especially, in arena of medical science and medicine, it is considered as next logical step and the future medicine [1, 2]. However, the potential risks such as harmful side effects on human and environment has made nanotechnology a double-edged sword [3,4].

Natural products based synthesis of nanoparticles can help to reduce the hazards of nanotechnology and thus can boost its applications. Biosynthesis, based on green chemistry principles, is proactive approach and has numerous advantages over conventional chemical and physical methods such as cost-effectiveness, environment friendly, simple and safe, no use of hazardous chemicals as reducing agents, less wastage of materials, minimum energy usage, safer disposal and recycling [5, 6]. Studies are also reported on the preparation of various types of nanostructures by using natural products such as vitamins, biodegradable polymers, enzymes, polysaccharides and different microorganisms including algae, fungi, bacteria and viruses. Nevertheless, plants extract-mediated synthesis of nanomaterials is considered advantageous due to stability of nanostructures, cost efficacy, availability of plants, lower contamination risks, require little maintenance, and simple to scale up as plants are nature’s ‘chemical factories’ [6,7].

Silver Nanoparticles are famous for their anti-bacterial, anti-dengue viral and anti-fungal activates [8]. AgNPs are also used in textile industry, food packing, cosmetics, paints and detergents owing to their anti-microbial activity [9]. Preparation of AgNPs can be carried out by using extract of different parts of plant such as roots, stem, leaves or fruit. In the synthesis of AgNPs by plants, polyphenols play the main role in degradation of different organics compounds. These plant extracts usually play the dual role in the synthesis process as reducing and capping agent. During green synthesis, the coatings of various biomolecules on surfaces of AgNPs not only improve their stability but also enhance their biocompatibility by reducing the toxicity risks [8, 9].

In this study, we choose *Azadirachta Azadarachta indica indica* (*A. indica*) and *Citrullus Colocynthis colocynthis* (*C. Colocynthis colocynthis*) for the synthesis of AgNPs because both of these plants are famous for good medicinal properties and exhibit effective biological activities. *A. indica* plant belongs to family of Meliaceae and has enormous uses in medical field from ancient times [10]. It is familiar as anti-bacterial, anti-fungal and anti-microbial plant as it has quercetin, ß-sitosterol and nimbinin in their leaves and azadirachtin in its seeds [11]. The *A. indica*...
plant extract has been reported to induce various biological activities including antiseptic, antifungal, antioxidant, anti-inflammatory, antimicrobial, and insecticidal activities [10, 11]. C. colocynthis-colocynthis (also known as bitter apple) is a member of Cucurbitaceae family. The C. colocynthis fruit contains many biomolecules including alkaloids, glycosides, flavonoids, and fatty acids which have are famous for important biological activities such as antimicrobial, antioxidants, cytotoxic, antidiabetic, antilipidemic, and insecticide [12]. That’s why it is considered one of the best medicinal plants and is used in different biomedical applications including anti-microbial, anticancer activities [12-14].

According to World Health Organization (WHO), mosquitoes can be listed as one of the deadliest organisms because millions of humans die due to different diseases caused spread by them. Mosquitoes are the potential vector for many deceases diseases like dengue, malaria, west nile, chikungunya, zika and yellow fever. Dengue fever incidence has increased 30-fold worldwide in the last 30 years which is caused by Aedes aegypti mosquito vector Dengue incidence has increased 30-fold worldwide in the last 30 years which is caused by dengue mosquito larvae; Aedes aegypti (A. aegypti) [15]. Treatment of these diseases is a challenge. To control the rapidly increasing mosquito-borne risks, researchers are working on different strategies. Since mosquitoes breed in water and at larval stage, it is easy to target them there. However, inhibiting larvae in water by conventional ways using pesticides can increase the toxic risks for environment and humans. Thus natural pesticides such as plant extracts can be simple and side effect free promising methodologies. The use of green chemistry based nanomaterials can further enhance the effectiveness and efficiency of such approaches [16-19].

In this work, A. indica and C. Colocynthis-colocynthis mediated AgNPs were manufactured synthesized, characterized and tested for their anti-dengue larvicidal activity against A. aegypti and it was found that these AgNPs can potentially be considered good anti-larvicidal agents.

2. Materials and Methods

2.1. Material and chemicals
Fresh leaves of A. indica were collected from the Botanical garden of University of the Punjab, Lahore, Pakistan and C. colocynthis fruit was obtained from local market of Lahore, Punjab, Pakistan. Both products were identified by the expert of the Department of Botany, University of the Punjab, Lahore. Late fourth instar larvae of A. aegypti were collected from local pond and identified by the expert of the Department of Zoology, Lahore College for Women University, Lahore Pakistan. Silver nitrate (AgNO₃) was of research grade and obtained from Merck (Germany). To prepare synthesis solutions, aqueous extracts and for all other purposes deionized (DI) water was used throughout the experiment.

2.2. Preparation of A. indica and C. colocynthis extracts
The aqueous extract of A. indica leaves was prepared following the method reported by Pragyan et al [20] with some modifications. To remove any dust and contaminations, first fresh leaves were rinsed carefully with running tap water then washed twice with deionized (DI) water and let them dry in air. 20 g of these leaves were taken and converted into very small pieces by cutting them which were transferred to 250 mL conical flask of 250 mlL boiled for 10 min in 100 mL deionized (DI) water. After boiling, the mixture was let to cool down cooled to room temperature filtered with Whatman No. 1 filter paper before storing at 4 °C for next step. Different steps of the preparation of A. indica leaves aqueous extract are shown in upper panel of figure 2. One can notice that greenish color of mixture solution (figure 1c, before boiling) changed to yellowish light brown color in final stage (figure 1d).

The aqueous extract of C. colocynthis fruit was prepared by the technique mentioned elsewhere [21] with few changes and whole process of extract preparation is demonstrated in upper panel of figure 2. Briefly, a greenish white C. colocynthis fruit (as shown in figure 2a) was cleaned by washing many times with fresh tap water to get rid of any debris or any other contaminations, then rinsed two times with DI water before cutting into small pieces. 50 g pieces of C. colocynthis were soaked in 200 mL of deionized (DI) water for 72 h at room temperature. Finally mixture was filtered 2 times with Whatman No. 1 filter paper to get greenish yellow color of the extract (figure 2c) and kept at 4 °C for further use.

2.3. Synthesis of extract-mediated silver nanoparticles
To prepare AgNPs using aqueous extracts, first precursor stock solution (100mM) was made by dissolving 1.69 g of silver nitrate (AgNO₃) into 100 mL of DI water.

A. indica extract-mediated AgNPs were prepared by the protocol of Aparajita et al. [22] with some amendments. Three different amounts of prepared extracts were used in our case, that is different amounts of as-prepared extract (250 μL, 500 μL and 1000 μL) were used for the same amount 20 mL of silver nitrate precursor solution (1mM) to obtain the optimized concentration. In figure 2-1 (upper middle panel), different phases of AgNPs preparation using the A. indica extract are presented. First of all, colorless precursor solution of AgNO₃ was heated to 70 °C with gentle magnetic stirring of at 200 rpm (figure 2a1e). Then A. indica extract was added dropwise into the solution and the mixture was kept for heating at 70 °C under continuous stirring for 10 min (figure 2b1f). The color of the solution changed with time from transparent to light yellow to yellowish brown indicating the formation of AgNPs (figure 2e1g). At the end, the solution was let to cool down cooled to room temperature and kept at -4 °C for further use. Depending upon the amount of the extract (reducing agents) the color of final stage sample (colloidal AgNPs) was found to change as can be noticed in lower panel of figure 2-1 (lower panel), from yellowish brown to reddish brown to reddish dark brown for 250 μL, 500
μL and 1000 μL of the extract were taken as optimized NPs based on Uv-Vis results and was used in larvicidal activity and other characterizations.

For the synthesis of AgNPs using aqueous extract of C. colocynthis, approach described by Shawkey et al. [23] was followed with some alterations. Briefly, 20 mL of 5 mM silver nitrate solution was mixed with different amounts (1 mL, 2 mL, and 4 mL) of C. colocynthis extract and then the mixture was heated to the boiling temperature (about 95 °C) under constant stirring at 200 rpm for 20 min. Finally, solution was cooled down to room temperature and store at -4 °C. Color changes during the synthesis process indicate different phases of the reduction reaction and are presented in figure 4-2. We synthesized AgNPs using three amounts of as-prepared extract of C. colocynthis viz. 1 mL, 2.5 mL, and 4 mL following aforementioned method to achieve the optimized concentration. All three colloidal AgNPs samples mediated by different amounts of extract with precursor and extract samples are shown in figure 4-2 (lower panel). Nevertheless, only 2.5 mL extract-mediated AgNPs were selected as optimized AgNPs based on Uv-Vis results and for larvicidal activity and other characterizations.

2.4. Characterization of synthesized silver nanoparticles

The above prepared AgNPs were characterized by different techniques. Optical properties of all colloidal samples were examined by measuring the absorbance spectra with the ultraviolet–visible (UV-Vis) spectroscopy (Shimadzu, UV-1800, Japan) in the wavelength range of 300-700 nm wavelength. The size, shape and elemental composition investigations of synthesized nanoparticles were conducted by scanning electron microscopy and energy-dispersive X-ray (Nova NanoSEM 450, USA). For SEM analysis, samples were prepared by using the drop casting methods to achieve sufficient AgNPs amount on the clean glass substrate. To avoid any probable charging effects, a very thin gold coating was deposited on the samples before SEM-conducting SEM analysis. The structural nature of prepared AgNPs was examined by X-ray diffractometer (JDX 3201M, Jeol, Japan). For XRD, again, samples were prepared by developing a thick enough film of each colloidal sample by drop casting methods on the glass substrate. To study the presence of different probable functional group on the surface of AgNPs, Fourier-Fourier transform infrared spectroscopy (IRTracer-100, Shimadzu, Japan) was conducted in the range of 500-3600 cm⁻¹.

3. Larvicidal activity tests

The larvicidal potential of both types of green AgNPs and their respective extracts at different concentrations against dengue larvae were studied according to the standard protocol recommended by WHO [24]. 25 fourth instar A. aegypti larvae collected from local pond were transferred into different beakers containing 200 mL deionized DI water. Five different concentrations of both types of AgNPs were prepared and tested in each case; for A. indica mediated AgNPs following concentrations were used: 1.25 mg/L, 2.5 mg/L, 5 mg/L, 10 mg/L, and 20 mg/L, and for C. colocynthis mediated AgNPs concentrations used were: (0.3 mg/L, 0.6 mg/L, 1.25 mg/L, 2.5 mg/L, and 5 mg/L). The test concentrations of both types of AgNPs were selected on the basis of pilot experiments in which LC₅₀ concentrations (1.25 mg/L and 0.3 mg/L for A. indica and C. colocynthis mediated AgNPs respectively) were found. In each beaker required amount of AgNPs were added to achieve to the predetermined concentration. In the case of extracts, three concentrations (1 mL, 2 mL, and 3 mL) of each type of extracts were chosen while only DI water was used as control. The experiment was conducted in triplicates under laboratory conditions (at 26 ± 2°C). All the beakers were covered with perforated aluminum foil for air circulation. During exposure experiment, larvae were not provided with any food. After 24 h of treatment period, the counting of alive and dead larvae was made to calculate percent mortality. Larvae which showed no motility after being disturbed with a needle were counted as dead. The data are shown as mean value and standard error of mean (Mean± s.e.; x) and Microsoft Excel program was used to make the graphs. The percentage mortality was determined by using formula given below

\[
\frac{\text{Number of dead larvae}}{\text{total number of larvae exposed}} \times 100
\]

4. Results and discussion

4.1. Synthesis mechanism of extract-mediated AgNPs

Figure 5-3 illustrates schematically the potential formation mechanism of AgNPs using the aqueous extracts of different biomaterials. First of all, the precursor (AgNO₃) is dissolved into the solvent (DI water) that provides Ag-ions (Ag⁺) into solution without producing any color. Addition of extract induces the reduction of these Ag-ions to free Ag-atoms (Ag₀) by gaining the electrons. This reduction can be termed as bio-reduction or plant-assisted reduction because it is caused by different biomolecules such as protein and phytochemicals present in the extract which serve as reducing agent [25]. This reduction reaction can be noticed by the color change of the solution which may be enhanced by heating. The color of the reaction solution may change from transparent to light yellow to yellowish brown and finally dark brown depending upon the type of the extract used (as shown in figures 3 and 4). This time dependent color change during the synthesis (10 min for A. indica AgNPs and 20 min for C. colocynthis AgNPs), actually, indicates the different phases of nucleation and growth process occurring in the reactions [26]. The free silver atoms (Ag₀) accumulate due to van der Waals interactions.
and Brownian motion to form Ag-nuclei (nucleation process). The growth process of these nuclei into AgNPs can occur in different ways including coalescence of these nuclei and Ostwald ripening of smaller NPs into bigger ones. However, both nucleation and growth process may occur simultaneously in reaction during the formation of AgNPs [27]. The biomolecules of the extracts also served as capping agents because no additional stabilizing agent was used. The final size of the synthesized AgNPs can be controlled by varying the concentrations of extracts and precursor solution [28].

4.2. UV-Vis spectroscopy analysis

UV-Vis spectroscopy is very common and effective way to study the formation of colloidal nanoparticles by their optical responses. In Metallic NPs such as AgNPs, the free movement of electron between the valance band and conduction bands is possible due to very narrow gap between them. These electrons on the surface of AgNPs produce the surface plasmon resonance (SPR) which is actually resonant oscillation of conduction electrons in response of the incident light. Due to the SPR absorption, nanoparticles in colloidal form produce different colors depending upon various factors e.g., size, shape and surrounding medium [25].

To confirm purity and to study the optical response of colloidal AgNPs, UV-Vis spectroscopy was conducted and obtained absorption spectra are displayed in figure 6a. The characteristics surface plasmon resonance (SPR) peaks of AgNPs using three different amounts of A. indica extract 250 µL, 500 µL and 1000 µL are shown in left panel (figure 6a1a). All three peaks were in the wavelength range of 412 nm to 416 nm indicating the pure silver nature of the particles. A slight red shift can be noticed with increasing concentration of the extract in the reaction solution. A similar trend of peak shifting towards the higher wavelength values (416 nm, 429 nm, 431 nm) by enhancing the amount of extract (1 ml, 2 ml and 4 ml) was also observed in the synthesis of C. colocynthis mediated AgNPs as illustrated in right panel (figure 6a4). This red shift might be due to the change in size of the AgNPs because, in green synthesis, extracts served as the reducing agents and variation in reducing agent concentration can affect the final size of the nanoparticles [29, 30]. Furthermore, occurring of only single absorption peak in each case indicates the formation of spherical shaped AgNPs [31].

4.3. Scanning electron microscope (SEM) analysis

SEM is the most prominent characterization technique to analyze shape, surface morphology, size distribution and elemental composition of nanomaterials. In order to examine the morphology of AgNPs prepared by both extracts, SEM analysis was conducted and the obtained micrographs with histograms of particle size distribution are presented in figure 25.

In case of AgNPs mediated from A. indica extract, small spherical shaped silver particles like bright tiny spots can be seen on the bigger bio-particles of the extract. At some places agglomeration of few smaller particles into small clusters can also be noticed (figure 24a5a). The smaller nanoparticles have very high surface energy and they agglomerate into larger sized particles to minimize their surface energy. There could be many reasons for agglomeration of these nanoparticles because these particles have very high surface energy and to reduce energy particles agglomerate and form larger sized particles [32]. The size distribution range of these spherical AgNPs was found to be from 12 nm to 24 nm as depicted in a magnified view of figure 5b and presented graphically in particle size histogram (figure 5c). The average particle size was measured to be $\pm 17\pm 4$ nm.

In the lower panel, SEM micrograph (figure 25a5d) reveals the size and morphology of AgNPs mediated from C. colocynthis extract. A nice random distribution of NPs over the bigger extract particles can be visualized. No cluster formation due to agglomerate of NPs was noticed. Again most of the NPs were found of spherical shapes with diameters in the range of 20 nm to 36 nm (figure 5e) and having average diameter of $26\pm 5$ nm, as shown by the particle size histogram (figure 5f).

To confirm the existence of elemental silver in the synthesized particles, Energy-dispersive X-ray (EDX) was conducted. EDX technique determines the elemental composition analysis on the basis of energy values of characteristics x-rays which are unique for every element. The obtained EDX results are presented in figure 66, in upper panel for A. indica extract-mediated AgNPs while in lower panel for C. colocynthis extract-mediated particles. Insets show the table of elements in each case.

In both cases, we can notice strong silver signal (37.08% for Ai-AgNPs and 38.5% for Ce-AgNPs) along with other signals. The intense optical absorption peak at around 3 KeV is considered a typical characteristic absorption band for metallic silver due to surface plasmon resonance [28]. Thus EDX established the presence of elemental silver in both types of extract-mediated NPs by the characteristic SPR absorption peaks which supported the UV-Vis results. The other signals such as C, O, Na, Mg, Au, Cl, and K in the case of by A. indica AgNPs and C, O, Na, Ca, Mg, Si, Au, Cl, Pd, and K (figure 6a6a) for the C. colocynthis AgNPs sample (figure 6b6b) can also be observed. The occurrence of weak gold (Au) signal can be attributed to thin gold coatings on the sample to avoid any charging. The Si peak may be due to presence of glass substrate. The appearance of other signals especially intense O and C peaks indicate the presence of other metabolites on the AgNPs surface due to the aqueous extract (A. indica leaves and C. colocynthis fruit). These metabolites play a vital role of capping and stabilizing agents in the biosynthesis of the AgNPs because they provide the stability to AgNPs by surrounding and developing a thin capping layer of organic molecules. This advantage of green synthesis approach not only reduce the cost but also minimize the toxicity caused by hazardous chemicals used for reducing and capping purposes in chemical methods [28,33,34].

4.4. X-Ray Diffraction (XRD) analysis

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For the structural analysis of materials, X-ray diffraction (XRD) is an eminent analytical technique. It is widely used as a primary analytical tool to investigate material purity and different structural parameters such as crystal structure, crystallite size, lattice parameter, crystal phase identification, and various crystal defects [25]. In figure 9, indexed XRD patterns of both types of extract–mediated AgNPs are exhibited.

One can notice six prominent peaks in the diffractogram of *A. indica* mediated AgNPs peaks (figure 9a). The peaks appearing at 2θ values of 38.5°, 46.5°, 64.7° and 77.1° can be assigned to diffraction planes (hkl) values of (111), (200), (220) and (311) respectively, according to COD ID No. 9013052 [35]. These distinct reflection planes confirmed the silver metallic nature with face centered cubic (FCC) crystalline structure of synthesized AgNPs. Moreover, two other unidentified intense peaks occurring at 2θ values of 28.1° and 32.4°, showing higher degree of crystallinity, can also be noted (labeled by stars). These peaks indicate the presence of some crystalline moieties or organic compounds deposited on surface of AgNPs from aqueous extract of *A. indica* leaves. Many other studies of AgNPs prepared by green synthesis using the plant extract have also mentioned the appearance of such additional peaks in their XRD spectra which are believed to occur due to crystalline impurities of extracts [36, 37].

In XRD pattern of AgNPs prepared by *C. colocynthis* as shown figure 9b, four prominent distinct diffraction peaks observed at 2θ value of 37.9° (111), 46.2° (200), 67.5° (220) and 77.1° (311) indicate the FCC crystal structure of resultant nanoparticle, according to COD ID No. 9013046 [38]. Again, some extra intense peaks at 27.8° and 32.2° were observed (star peaks) which may be attributed to existence of crystalline nature of biomolecules on the AgNPs surface due to *C. colocynthis* extract [39].

To determine the crystallite size (D) of both type of extract–mediated AgNPs, the width of prominent Bragg’s reflection (111) in each case was utilized in the Debye–Scherer formula [26].

\[
D = \frac{k \lambda}{\beta \cos \theta}
\]

In this equation (4.1), \(k\), \(\beta\), \(\theta\) and \(\lambda\) are x-ray wavelength (1.54 Å), shape factor with value of 0.9, peak width (full width half maximum–FWHM) and Bragg diffraction angle respectively. The average particle-crystallite size determined by XRD for AgNPs prepared using extracts of *A. indica* and *C. colocynthis* were found to be 11±1 nm and 15±1nm respectively (table 1) which has fair agreement with particle size measurements obtained by SEM.

| Sample                  | Peak position (20) | Diffraction plane (hkl) | FWHM (rad) | Crystallite size, D (nm) | Lattice parameter, a (nm) |
|-------------------------|-------------------|-------------------------|------------|--------------------------|--------------------------|
| *A. indica* extract–mediated AgNPs | 38.5°             | (111)                   | 0.0128     | 11±1                      | 0.403                    |
| *C. colocynthis* Extract–mediated AgNPs | 37.9°             | (111)                   | 0.0095     | 15±1                      | 0.409                    |

### 4.5. Fourier-Transform Infrared Spectroscopy (FTIR) Analysis

FTIR spectroscopy is a very useful and powerful technique to identify different functional groups and chemical bonds in any sample. FTIR characterizes the chemical composition of the material by interaction of infrared radiation with a surface of the sample. The infrared radiations are absorbed by the sample at specific frequency ranges depending upon the type of chemical functional groups and bonds. The measurement of these frequencies and intensities help to study the chemistry of the sample.

In this work, FTIR analysis was carried out and obtained spectra are presented in figure 8. The aim of FTIR study was to examine the presence of different biomolecules and functional groups deposited on the surface of both types of AgNPs. These biomolecules helped in bio-reduction and efficient stabilization of NPs during the synthesis process. In this work, to study the presence of different biomolecules and functional groups on the surface of AgNPs, in both cases, which provided the required bio-reduction and efficient stabilization during the synthesis process, FTIR measurement were carried out and obtained spectra are presented in figure 10.

FTIR spectrum of AgNPs synthesized from *A. indica* leaves is shown in figure 10a where various absorption bands at different specific frequency indicate a composite nature. The prominent peaks occurred at the frequency ranges of 3265, 2919, 2849, 2363, 1734, 1616, 1244, 1161, 1097 and 1027 cm⁻¹ as can be noticed from the figure 10a. A broad band appearing in the range of 3265 cm⁻¹ can be attributed to the stretching vibrations of O-H bond indicating the presence of polyphenols. The peaks occurring at value range of 2919 cm⁻¹ and 2849 cm⁻¹ can be assigned to C=C aromatic and C-H alkaline (asymmetric stretching of C-H bonds) groups [40]. The possible presence of C=N and C≡C triple bonds can be ascribed by peaks of 2363 cm⁻¹ because band appearing in the range of 2200-2400 cm⁻¹ indicate the C-N and C≡C triples bonds. The band at 1734 cm⁻¹ could be due to C=O stretching vibration in the carbonyl groups from terpenoids and flavonoids. The peaks of 1616 cm⁻¹ might indicate presence of C=C bonds or aromatic rings attributed to carbonyl stretch in proteins [41]. The absorption bands appearing at 1244, 1161, 1097 and 1027 cm⁻¹ can be
assigned to the C-H alkene (Aliphatic aldehydes, C-H asymmetrical stretching functional group) and C-O vibrations of ether linkages [14, 40, 42]. Thus FTIR analysis of *A. indica* mediated AgNPs confirmed the presence of different phytochemical components such as flavonoids, polyphenols and terpenoids of *A. indica* extract which potentially played a vital role in the reduction of Ag⁺ ions to Ag⁰ atoms. Especially the proteins (due to the carbonyl and NH, of amino acid) present in *A. indica* leaves extract served as reducing and encapsulating agent because carbonyl group can reside on the metallic silver due to strong affinity for metals and thus developing a capping layer on surfaces of AgNPs [20, 36, 40, 42].

For the AgNPs prepared by *C. colocynthis*, FTIR spectrum exhibited peaks at 3255, 2920, 2341, 1582, 1240, 1141, 1095, 1007 and 953 cm⁻¹ as shown in figure 10b. The occurrence of absorption bands at 953-1007 cm⁻¹ might be due to C-O- or C-O-C- functional groups while the peaks at 1010-1150 cm⁻¹ can be allocated to C-N stretching vibrations related to aliphatic amines or to alcohols or phenols indicating the existence of polyphenols [14, 43]. The bands at 1240 cm⁻¹ and 1339-1460 cm⁻¹ attributed to the amide III and II groups respectively. Likewise the absorption in this region (around 1384 cm⁻¹) indicated the residual amount of NO₃. Whereas the prominent absorption band at 1582 cm⁻¹ might be owing to symmetric stretching vibrations of -COO- groups of amino acid confirming the presence of *C. colocynthis* *A. indica* extract protein. The existence of proteins was further indicated by the peak at 1634 cm⁻¹ due to stretch vibration of -C=O of amide I bonds of proteins. The bands arising at around 2200-2400 cm⁻¹ indicate the probable presence of C-N or C-C triple bonds. The peaks occurring 2920 cm⁻¹ can be dispensed to C-H bond vibrations. The intense broad absorption band at 3355 cm⁻¹ can be designated to characteristic of –OH (hydroxy) group indicating the existence of alcohols and phenolic compounds [34, 44, 45]. The FTIR results confirmed that the *C. colocynthis* mediated AgNPs were surrounded by various phytochemical constituents such as amines, aldehydes, alcohols, ketones and carboxylic acids. These *C. colocynthis* extract based biomolecules (proteins and metabolites) played dual role of bio-reduction and stabilization during the synthesis of AgNPs.

### 4.6. Larvicidal Bioassay

In order to study the antilarval activity of aqueous extracts of *A. indica* and *C. colocynthis* and their respective AgNPs, different concentrations were tested for larvicidal bioassay against fourth instars larvae of *A. aegypti*. In order to study the antilarval activity of prepared samples; aqueous extracts of *A. indica* and *C. colocynthis* and AgNPs mediated by these extracts at different concentrations, were subjected separately to larvicidal bioassay on fourth instars larvae of dengue (*A. aegypti*). After 24 h treatment, dead and alive mosquito larvae were counted to calculate percent mortality. Larvae which showed no motility after being disturbed with a needle were considered dead. The typical demonstration of larvicidal bioassay are exhibited in figure 12 (A. indica) and figure 12 (D. colocynthis) panel).

For *A. indica* leaves aqueous extract, three concentrations; 1 ml, 2 ml and 3 ml, 0.5 %, 1 % and 1.5 % of as-prepared extract was tested while five concentrations 1.25 mg/L ppm, 2.5 mg/L ppm, 5 mg/L ppm, 10 mg/L ppm and 20 mg/L ppm of *A. indica* extract mediated AgNPs were exposed to *A. aegypti*. DI water was taken as control in each experiment. In the initial stage of this bioassay at t= 0 h, (as shown in upper panel of figure 12), random scattering of the larvae indicate their active movement in all the samples. After 24 h, lower panel of figure 12, all larvae in the control (water) were alive and moved energetically. Beakers containing extract of *A. indica* showed no mortality in all three concentration samples. Nonetheless larvae were found less motile in the sample with 3 mL concentration of extract as compared to other two samples (1 mL and 2 mL extract). Beakers containing AgNPs showed different mortalities with different concentrations. The death larvae can be observed at the bottom or gathered in the middle of each beaker with high concentration (lower panel of figure 12). AgNPs with concentration of 1.25 mg/L ppm, 2.5 mg/L ppm, 5 mg/L ppm, 10 mg/L ppm and 20 mg/L ppm caused 59%, 66%, 80% and 85% and 100% mortality respectively. All the larvicidal results of *A. indica* leaves extract and mediated AgNPs are summarized in table 1.

**Table 2** Larvicidal activity of DI water (control), aqueous extract of *A. indica* leaves and extract-mediated AgNPs at different concentrations against fourth instars larvae of *A. aegypti*.

| Sample type          | Concentration of sample | Percentage Mortality (%) |
|----------------------|-------------------------|--------------------------|
| Water (control)      | -                       | 0 (active)                |
| as-prepared extract  | 1 ml (0.5 %)            | 0 (active)                |
| A. indica leaves     | 2 ml (1 %)              | 0 (active)                |
| A. indica extract-     | 1.25 mg/L ppm          | 49 (active)               |
| mediated AgNPs       | 2.5 mg/L ppm           | 66 (active)               |
|                      | 5 mg/L ppm             | 80 (active)               |
|                      | 10 mg/L ppm            | 85 (active)               |
|                      | 20 mg/L ppm            | 100 (active)              |

A similar larvicidal activity assay was conducted for samples of *C. colocynthis* fruit extract; in this case again three extract concentration of as-prepared aqueous extract; 1 ml, 2 ml and 3 ml, 0.5 %, 1 % and 1.5 % and DI water as control were taken while for extracted mediated AgNPs five concentration were chosen as 0.3 mg/L ppm, 0.6 mg/L ppm, 1.25 mg/L ppm, 2.5 mg/L ppm and 5 mg/L ppm (figure 12). In the beginning of the activity test (at 0 h), all larvae were alive and motive in samples (upper panel of figure 12). After exposure of 24 h, again no mortality in (control) and all three extract samples, however, a concentration dependent effect on motility of larvae was noticed in extract samples. The
larvae in sample with 0.3 ml2% extract. In contrast, about 0.1 ml1.5% extracts, highest mortality was counted in the sample with maximum concentration. A mortality of 53%, 64%, 80%, 91% and 100% were recorded for samples with AgNPs concentration of 0.3 mg/L ppm, 0.6 mg/L ppm, 1.25 mg/L ppm, 2.5 mg/L ppm and 5 mg/L ppm respectively. Assembly of immotile (dead) larvae can be witnessed in middle of beakers (lower panel, figure 1210). The statistics of C. colocynthis mediated larvicidal assay are listed in table 23.

Table 3 Larvicidal results of DI water (control), aqueous extract of C. colocynthis fruit and extract-mediated AgNPs at different concentrations against fourth instar larvae of A. aegypti. 

| Sample Type                  | Concentration of sample | Percentage Mortality (%) |
|------------------------------|-------------------------|--------------------------|
| Water                        | -                       | 0 (active)               |
| as-prepared aqueous extract of C. colocynthis fruit | 1 ml0.5% | 0 (active) |
|                              | 2 ml1%                  | 0 (less motile)          |
|                              | 3 ml1.5%                | 0 (least motile)         |
| C. colocynthis extract-mediated AgNPs | 0.3 mg/L ppm | 53±4 |
|                              | 0.6 mg/L ppm            | 64±6                     |
|                              | 1.25 mg/L ppm           | 80±3                     |
|                              | 2.5 mg/L ppm            | 91±3                     |
|                              | 5 ppm/mg/L              | 100±0                    |

We did not observe any larvicidal efficacy effect of both types of aqueous extracts (A. indica leaves and C. colocynthis fruit) was noticed very insignificant as compared to AgNPs mediated by these extracts. Some other researchers have also reported negligible biocidal activity of aqueous extracts in comparison to extracts mediated nanoparticles [21, 23, 47]. On other hands, a concentration-dependent biocidal efficiency of pure aqueous extracts of various natural products can also be found in literature [16, 28]. It seems that biocidal activities of extracts depend not only on the concentrations used in the bioassays but also on the level of purification (crude, highly purified, aqueous, or alcoholic) as well as nature and different parts of bio-materials to be extracted such as leave, fruit, stem and seed [16, 19, 46].

In the case of extract-mediated AgNPs, the enhanced larvicidal activity of AgNPs can be attributed to the synergistic effect of Ag-ions and deposited biomolecules. The difference in the larvicidal efficiency occurs due to the presence of different types of biomolecules on the surfaces of AgNPs though Ag-ions were same in both cases. In both types of extract mediated AgNPs, We observed, in both cases, that by increasing the concentration of AgNPs, percent mortality of larvae increased. Nevertheless, C. colocynthis fruit extract mediated AgNPs were found more effective at lower concentrations and exhibited strong larvicidal efficacy as compared to A. Indica leaves extract mediated AgNPs as demonstrated graphically in figure 13. As mentioned above, different types of biomaterials (plants) and even various parts of same plant may affect the biocidal efficiency of mediated AgNPs. That’s why perhaps AgNPs prepared by fruit extract of C. colocynthis showed better antilarval activity even at lower concentrations in our case. A comparative larvicidal study of extracts and mediated AgNPs using various parts such as leaves, fruit, seed and stem of same plant can be conducted for more detailed investigations.

Since the exact mechanism of larvae mortality is unidentified, different modes of action can be proposed for larvicidal activity of AgNPs. Most probably the size and surface chemistry of AgNPs play a vital role to make them lethal against mosquito larvae. At the first stage, owing to small size AgNPs penetrate into the larval membrane causing the possible leakage of cellular materials. In the next step, AgNPs may interact with the different cell molecules to inhibit molting and other physiological processes. In their action, AgNPs may produce peroxide, inactivate the enzymes and disturb the functional proteins leading to the larva death. The presence of extract biomolecules on the AgNPs surface can further enhance their effectiveness in this larvicidal process [23, 40, 46, 47]. This indicates that anti-dengue antilarval pharmaceuticals can be formulated using very low concentration of natural products based AgNPs for practical applications. Thus findings of this study can be helpful to provide new paradigm in designing the green chemistry based alternative to combat the increasing mosquitoes diseases.

5. Conclusion

AgNPs were prepared from herbal origin which is economical and eco-friendly using the aqueous extracts of A. indica leaves and C. colocynthis fruit. Both types of NPs were found pure metallic silver in nature having FCC crystalline structure and were spherical in shape with diameters of 25±5 nm, as determined by SEM and XRD. The optical behavior of colloidal samples and presence of different extract biomolecules were studied by UV-Vis and FTIR.

Both types of NPs were found pure metallic silver in nature having FCC crystalline structure and of spherical in shapes with diameters in the range of 25±5 nm, synthesized by different biomolecules of extracts as determined by different characterization techniques including SEM, UV-Vis, XRD and FTIR. Both types of The prepared green AgNPs exhibited different great larvicidal potential against A. aegypti even at low concentrations; LC50 at 1.25 mg/L ppm for A. indica AgNPs and LC50 at 0.3 mg/L ppm for C. colocynthis mediated AgNPs respectively. We observed a concentration dependent larvicidal activity in both types of AgNPs whereas their extracts showed negligible efficiency against larvae in both cases. These results indicate the potential use of green AgNPs as alternative to replace the synthetic products in pharmaceuticals for anti-dengue antilarval purposes.
Acknowledgments
Authors acknowledge Dr. Shahid Atiq and Dr. Syed Sajjad Hussain from Centre of Excellence in Solid State Physics, University of the Punjab, Lahore-Pakistan for their cooperation in experimental work and useful discussions in manuscript writing.

Ethical Statement
It is not relevant to our work.

Funding Statement
This work was financially supported by Higher Education Commission (HEC) of Pakistan under the Project of ‘National Research Program for Universities’ (Project No.: HEC-NRPU-8019).

Data Accessibility
Our data are deposited at Dryad: https://datadryad.org/stash/dataset/doi:10.5061/dryad.z612j6m68k [48], with following reviewer URL; Dryad Reviewer URL: https://datadryad.org/stash/share/DksW82FCDqISwPqI2aYhDl4C9QUE74hEBPTmrJ3trHw

Competing Interests
We have no competing interests.

Authors’ Contributions

1). SR and MAR performed the synthesis experiments, analysed the data and wrote the manuscript
2). ZK, SR and MAR designed the experiments and conducted the larvicidal bioassays
3). SR and SN conducted the characterization measurements and helped in interpretation of data
4). FM and ZK provided the larvae facility and helped in larvicidal activity experiments
5). SR and SN conducted the characterization measurements and helped in interpretation of data
6). MJR and SR conducted the FTIR and SEM characterization and helped in data analysis

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Figure Captions

**Figure 1.** Upper panel-Different steps during the preparation of aqueous extract of *A. indica* leaves; Middle Panel- Synthesis of AgNPs; Lower Panel-(h) precursor solution, (i) reducing agent, (j-l) colloidal samples of AgNPs prepared by different amounts of *A.indica* extract.

**Figure 2.** Upper panel-Different steps during the preparation of aqueous extract of *C. colocynthis* leaves; Middle Panel- Synthesis of AgNPs; Lower Panel-(h) precursor solution, (i) reducing agent, (j-l) colloidal samples of AgNPs prepared by different amounts of *C. colocynthis* extract.

**Figure 3.** Schematic illustration of the synthesis mechanism of extract-mediated AgNPs. The phenomena occurring in each phase is mentioned accordingly.

**Figure 4.** UV-Vis spectroscopy results of AgNPs synthesized by (a) three different concentrations of *A. indica* leaves extract and (b) three different concentrations of *C. colocynthis* fruit extract. Single absorption peak in each case appeared in the wavelength range of 412-430 nm indicating the spherical shaped AgNPs. Insets display images of each colloidal sample in cuvette.

**Figure 5.** SEM micrographs and histograms. (a,b) SEM image of *A. indica* AgNPs showed spherical shaped NPs decorated on larger extract particles, (c) Histogram of *A. indica* AgNPs indicated their diameter ranging between 12-24 nm, (d,e) SEM results of *C. colocynthis* AgNPs reveal the sphere like morphology and random dispersal over the bigger extract particles. (f) Histogram of *C. colocynthis* AgNPs indicated their diameter ranging between 20-36 nm.

**Figure 6.** EDX spectrum showing elemental composition analysis of AgNPs (a) *A. indica* leaves, (b) *C. colocynthis* fruit. Insets show the table of elements in each case.

**Figure 7.** X-ray diffraction patterns of (a) *A. indica* AgNPs and (b) *C. colocynthis* AgNPs.

**Figure 8.** FTIR spectra (a) *A. indica* AgNPs, (b) *C. colocynthis* AgNPs. Occurrence of various absorption peaks at different positions indicates the presence of different biomolecules on the surface of NPs.

**Figure 9.** A representative display of larvicidal bioassay. Water (control), *A. indica* extract (Conc: 1.5 %), AgNPs at different concentrations. Upper panel shows the initial stage of experiment with 0 h time. Lower panel shows results after 24 h.

**Figure 10.** A typical demonstration of larvicidal activity. Water (control), *C. colocynthis* extract (Conc: 1.5 %), AgNPs at different concentrations. Upper panel shows stage in the start of the experiment (at 0 h). Lower panel displays the final stage story (after 24 h).

**Figure 11.** Graphs showing the larvicidal effect of AgNPs at different concentrations (a) *A. indica* AgNPs showed LC$_{50}$ at about 1.25 mg/L while at 20 mg/L 100% mortality was observed, (b) *C. colocynthis* AgNPs exhibited LC$_{50}$ at 0.3 mg/L and 100% mortality was observed at 5 mg/L.

**Figures**
**Figure 1**

- (a) Fresh *A. indica* leaves
- (b) Leaves after drying
- (c) Boiling process
- (d) Extract after filtration
- (e) Aqueous AgNO$_3$ solution during heating
- (f) Dropwise addition of extract during heating and stirring
- (g) Final stage of *A. indica* extract mediated AgNPs formation
- (h) Aqueous solution of AgNO$_3$
- (i) Aqueous extract of *A. Indica*
- (j) 1000 µL extract mediated AgNPs
- (k) 500 µL extract mediated AgNPs
- (l) 250 µL extract mediated AgNPs
Figure 2

(a) Green C. colocynthis fruit
(b) Small pieces soaked in water
(c) Extract after filtration
(d) Aqueous precursor (AgNO₃) solution
(e) Heating and stirring after mixing extract
(f) Color change during reaction process
(g) Final stage of C. colocynthis extract mediated AgNPs formation
(h) Aqueous solution of AgNO₃
(i) Aqueous extract of A. Indica
(j) 4 mL extract mediated AgNPs
(k) 2 mL extract mediated AgNPs
(l) 1 mL extract mediated AgNPs

Figure 53

Silver nitrate (AgNO₃) → Creation of silver ions (Ag⁺) → Reduction process Ag⁺ → Ag⁰ → Formation of silver nuclei

1. Stirring and heating
2. Stirring and heating
3. Stirring and heating
4. Stirring and heating

Nucleation and Growth processes to develop AgNPs
**Figure 64**

**Figure 75**
Figure 86

Figure 97
Figure 108

(a) A. indica mediated AgNPs

(b) C. colocynthis mediated AgNPs

20 mg/L AgNPs 10 mg/L AgNPs 5 mg/L AgNPs 2.5 mg/L AgNPs 1.25 mg/L AgNPs Extract (A. indica) Water

Figure 119

5 mg/L AgNPs 2.5 mg/L AgNPs 1.25 mg/L AgNPs 0.6 mg/L AgNPs 0.3 mg/L AgNPs Extract (C. colocynthis) Water

Figure 1210
Figure 1311

(a) *A. indica* AgNPs

(b) *C. colocynthis* AgNPs

[Bar charts showing varying mortalities at different concentrations for *A. indica* and *C. colocynthis* AgNPs.]
We are thankful to the reviewers for their expert assessment of our manuscript. We are grateful for their thoughtful and valuable comments to improve our manuscript. All points and issues raised by the learnt reviewers have been considered and the manuscript has been modified accordingly. A detailed point-by-point response to all comments is provided below indicating the implemented changes in the revised version of the manuscript. A highlighted version by ‘Track Changes’ is also included with the resubmission.

Response to 1st Reviewer’s Comments

Comment 1. The article describes the synthesis of Ag nanoparticles using *Azadirachta indica* and *Citrullus colocynthis* extracts. The work done is useful and is in line with the current scientific demands. There are also some issues to be addressed before publishing this work.

Response 1. We are grateful to the learnt reviewer for mentioning our work “useful “and “in line with current scientific demands”.

Comment 2. There are several grammatical mistakes for instance I am highlighting some of these:

i) page 2 line 17 which have famous is incorrect it should be which are famous

ii) page 2 line 25 “vector for many deceases like” it should be vector for many diseases like

iii) page 2 line 25 please correct “fever.Dengue”

iv) page 2 line 35 “AgNPs were manufactured” were synthesized

v) page 2 line 35 “the mixture was let to cool down to room temperature” should be the mixture was cooled to room temperature

The authors should thoroughly check the whole manuscript and English should be improved as well as all the editorial mistakes should be rectified.
Response 2. We are thankful to the reviewer for pointing out these grammatical mistakes to improve beauty of the text. The whole manuscript has been thoroughly checked and all the grammatical and editorial mistakes mentioned by the reviewers and otherwise are corrected.

Comment 3. Page 6 line 10, authors mentioned the particle size, I guess it should be crystallite size?

Response 3. The corrections were made as suggested by the reviewer.

Comment 4. Authors have not mapped the particle size in SEM images. The particle size should be measured in SEM image and it should be visible in the picture.

Response 4. According to the reviewer recommendations, the SEM images have been modified by adding the new images showing particle size visible in the picture.

Comment 5. The authors have mentioned that the chosen plants have known biological activity. However, they do not show any activity against A. aegypti. Secondly, if the extracts are not active against A. aegypti why there is a large difference in activity of AgNPs prepared from two different extracts as LC50 at 1.25 ppm for A. indica and LC50 at 0.3 ppm for C. colocynthis is observed. The later is 4 times more active than the former one. I do not seem that activity is only due to AgNPs in that case it should be similar or close to each other.

Response 5. We are appreciative to the reviewer for highlighting this point. Firstly, it is well-established fact that various medician plants including *A. indica* and *C. colocynthis*, exhibit different biological activities (R1-R3). However, in our case, both of the aqueous extracts (*A. indica* leaves and *C. colocynthis* fruit) did not show any larvicidal efficacy against dengue vector. The inefficiency of aqueous extracts of different plants was also reported by other researchers [R3-R6] however, some studies report concentration-dependent biological efficacy of aqueous extracts of some natural products [R2,R7]. Thus, biological activity of extracts of biomaterials depend on various
factors such as nature, different parts (leave, fruit, stem and seed) and level of purification (crude, highly purified, aqueous, or alcoholic) and concentration [R8-R9].

The reason of the difference in larvicidal activity of AgNPs prepared by *A. indica* leaves or *C. colocynthis* fruit is that different types of biomolecules are deposited on the surface of AgNPs (as confirmed by FTIR analysis). This is, in fact, the synergistic effect of Ag-ions and biomolecules to enhance the larvicidal performance of extract mediated AgNPs. Thus the difference in larvicidal efficiency occurs due to the presence of different types of biomolecules on the surfaces of AgNPs although Ag-ions are same in both cases. Therefore, the LC$_{50}$ values for both cases were different (LC$_{50}$ at 1.25 ppm for *A. indica* and LC$_{50}$ at 0.3 ppm for *C. colocynthis*).

Main text has also been modified to explain this point.

**Comment 6.** Figures 1, 2 and 5 are unnecessary and should be deleted.

**Response 6.** Figures 1 and 3 and figures 2 and 4 are combined according to the recommendations of reviewer 2, instead of deleting.

Figure 5 depicts schematics of the potential formation mechanism of extract mediated AgNPs and different phases of nucleation and growth processes. We would like to keep it in the main text with the kind consent of our proficient reviewer.

Data of all modified figures has been uploaded to Dryad Digital Repository

Our data are deposited at Dryad:

https://datadryad.org/stash/dataset/doi:10.5061/dryad.z612jm68k

Dryad Reviewer URL:

https://datadryad.org/stash/share/DksW82FCDqISwPql2aYhDi4C9QUE74hEBPTmrJ3trHw

**Comment 7.** The sentence in conclusion section “Both types of NPs were found pure metallic silver in nature having FCC crystalline structure and of spherical in shapes with diameters in the range of 25±5 nm, synthesized by different biomolecules of extracts as determined by different characterization techniques including SEM, UV-Vis, XRD and FTIR” is too long and understandable. It should be refabricated to two sentences.

After addressing these issues the article can be published
Response 7. We are agreed with the reviewer and this sentence has been rephrased into two sentences as,

“Both types of NPs were found pure metallic silver in nature having FCC crystalline structure and were spherical in shapes with diameters in the range of 25±5 nm, as determined by SEM and XRD. The optical behavior of colloidal samples and presence of different extract biomolecules were studied by UV-Vis, and FTIR.” The main text has also been modified accordingly

We are obliged to the reviewer for recomemnding our manuscript for “publication” in RSOS.

Response to 2nd Reviewer’s Comments

Comment 1. The MS presents work which has been published with various other plants. The work carried out is significant but need major revision.

Response 1. We are obliged to the reviewer for mentioning our work “significant”.

Comment 2. MS needs enormous revision in scientific reporting, language and grammar. The scientific names are wrongly spelled at some places.

Response 2. We are grateful to the learnt reviewer for highlighting the errors to enhance the flouncy and beauty of the text. The manuscript is carefully examined and all mistakes of language, grammar, typos and scientific reporting including scientific names pointed out by the reviewer and otherwise are corrected and highlighted by ‘Track Changes’ in the revised manuscript.

Comment 3. Each section of the MS needs major revision. The comments have been marked on the MS. Authors are advised to go through each and every comment; and rectify accordingly.

Response 3. We are thankful to the reviewer for valuable comments and suggestions. We have addressed all the comments and points raised by the learnt reviewer and
manuscript has been thoroughly revised in the light of remarks and recommendations of the reviewer.
The detailed response to the comments is described below to highlight the amendments made in the revised version of the manuscript accordingly.

**Comment:** The MS is about anti-dengue vector. Kindly revise.

**Response:** The title of the MS has been rectified as directed by the reviewer.

**Comment:** Check series.... FTIR was done in last..Please write based on conducting the characterization

**Response:** In abstract, series of the characterizations techniques has been corrected, as suggested by the reviewer.

**Comment:** Authors have not added any comparison between the characteristics of two kinds of NPs. Some features may be mentioned.

**Response:** We agreed to the reviewer and the abstract has been modified by adding comparative characteristic features of both types of AgNPs.

**Comment:** write scientific names correctly throughout the manuscript. Start specific names with small case. Take care of spellings. A. indica and C. colocynthis

**Response:** All scientific names and terms have been corrected including A. indica and C. colocynthis, as directed by the reviewer.

**Comment:** Which part? As you have prepared NPs with fruits, please specify

**Response:** The C. colocynthis fruit has been specified and reference has also been added in the main text, as suggested by the reviewer.

**Comment:** Dengue is transmitted by Aedes mosquitoes. It is caused by virus. Larvae have no role in it. Please rectify Write as hyphenated throughout the MS
Response: We agreed to the reviewer and the sentence has been rectified in the main text as; “Dengue fever incidence has increased 30-fold worldwide in the last 30 years which is caused by dengue virus”

Comment: What do you mean by glass substrate?
Response: Glass substrate means just a small piece of glass slide on which few drops of colloidal AgNPs sample are dried to make a thick enough NPs layer. This was used as sample for characterization purposes. It has also been described in literature [R10, R11].

Comment: 1ml, 2.5 ml and 4ml……………….. on what basis the ratios (pattern of conc.) were decided? The amount/conc. of a compound to be used in a mixture is always in a particular ratio. For example…1, 2, 4, 8….

However, only AgNPs prepared by using the 500 µl of extract was used in larvicidal activity and other characterization, specify the reason plz

Response: In this study, two types of extracts were used as reducing and stabilizing agents for synthesis of AgNP. The amount of extracts in both cases was selected from the literature protocols [R12, R13]. To obtain the optimized properties three different amounts of each extract were used for the preparation of AgNPs. An amount was selected and then its 2 fold and 4 fold was used for AgNPs synthesis process. In the case of A. indica extract, 250 µL, 500 µL and 1000µL while for C. colocynthis extract 1 mL, 2 mL and 4 mL were used (2.5 mL was typo error which has been corrected in the main text). The AgNPs prepared from 500 µL (A. indica) and 2 mL (C. colocynthis) were taken as optimized AgNPs on the basis of Uv-Vis results. This point has been addressed in the main text according to the reviewer’s recommendations.

Comment: Why did authors use directly 5 mM concentration to prepare NPs? Higher concentration of silver nitrate itself is toxic? Did you try for lower concentrations? What was the result? If not, then why?
Response: We agreed to the reviewer that AgNPs can be made by different concentrations of precursor AgNO₃ solution (Lower and higher than 5mM). In this study, both types of AgNPs were prepared following the protocols mentioned in the literature. A. indica extract-mediated AgNPs were synthesized using 1mM solution of AgNO₃ using methods described by of Aparajita et al. [R12] while C. colocynthis extract-mediated AgNPs using 5mM concentration of precursor solution as reported by Shawkey et al. [R13].

Comment: We cannot say that the fruit extract was more effective at lower concentration as the comparison is between 1 mM and 5 mM of AgNO₃. 5 mM is itself a very high concentration of silver nitrate. If authors want to compare the efficacy then they should do at similar concentrations either 1 mM or 5 mM.

Response: In our work we used two different types of extracts and AgNPs to study the larvicidal activity. Each type of AgNPs were prepared following different protocols using different precursor concentration (1mM or 5mM) and two different reducing agents (A. indica leaves extract or C. colocynthis fruit extract). We aim to report the antilarvae performance of each type of extract and AgNPs at different concentrations. It is well-known fact that change in concentration of precursor solution will affect the size and morphology of prepared AgNPs [R14, R15]. In our case, we used two type of extracts as reducing and stabilizing agents and thus different types of biomolecules were deposited on the surface of each type of AgNPs which caused variation in larvicidal efficacy.

We have rectified the main text (abstract, result and discussion and conclusions) to address the comparison issue raised by the reviewers.

Comment: ppm unit is no longer in use. May be replaced.

Response: According to reviewer recommendations, ppm has been replaced by mg/L throughout the manuscript.

Comment: “The test concentrations of both types of AgNPs were selected on the basis of pilot experiments in which LC50
concentrations were found”………..Any published result? If so, give ref. Otherwise, you may give values obtained here

Response: The LC$_{50}$ values of pilot experiments were found to be 1.25 mg/L and 0.3 mg/L for *A. indica* and *C. colocynthis* mediated AgNPs respectively. These LC$_{50}$ values have also been mentioned in the main text according to the reviewer suggestions.

Comment: This time dependent color change during the synthesis, actually, …………time not mentioned

Response: Time has been mentioned in the main text as “10 min for *A. indica* mediated AgNPs and 20 min for *C. colocynthis* mediated AgNPs.”

Comment: we can notice strong silver signal along with some other signals. mention Ag%

Response: % age values of Ag have been mentioned for both types of AgNPs in main text as “37.08 % for *A. indica* AgNPs and 38.5% for *C. colocynthis* AgNPs” as directed by the reviewer.

Comment: mL denotes the volume. As per WHO protocol, 1 mL of toxicant is added to 249 mL of water to study larval mortality. What about concentration? Write the conc of extract. How can you compare the mL of extract with ppm of AgNPs?

Response: The reviewer is right that ml determines the volume only. We have added percentage concentration of the extract. It is in common practice to use the extract in percentage concentration [R16, R17]. The prepared extract is standardized as 100 % from which dilutions are drawn for biassays. We used 3 different amounts (1mL, 2mL and 3mL) of the extract to be dissolved in water to make a final volume of 200 mL. 1 mL of extract in 200 mL of water will make a 0.5 % extract concentration, 2 mL will make 1 % and 3 mL will make 1.5 % extract concentration in the solution. Furthermore because the dynamics of the concentration formulations for extract and nanoparticles is different, extract is based on the extracted material while nanoparticles’ concentration is
devised from the precursor used, therefore, it is not necessary to have a direct comparison of the two in terms of concentration units.

**Comment:** 1ml, 2ml, 4 ml............ Are these concentrations? , 2, 3 mL..... Is this concentration?

**Response:** No, these are quantities, now we have added % age concentration of extracts (0.5 %, 1%, and 1.5%) in main text to rectify this issue raised by the reviewer.

**Comment:** How can you make anti-dengue medicine which has been tested against dengue larvae not dengue virus

**Response:** We are thankful to the reviewer for indicating this point, we have rectified the main main text as “This indicates that antilarval pharmaceuticals can be formulated using very low concentration of natural products based AgNPs for practical applications”

**Comment:** Figure captions: all figure captions need to be revised...as they too lengthy, unnecessarily too big legends. Please make all legends concise and crisp.

**Response:** All the figure captions have been revised to make brief and concise, as directed by reviewer.

**Comment:** Figs. 1, 2 and 3 can be easily combined. Please shorten and make it crisper. F1: explanations can be included in the figures itself (labellings of figures)

**Response:** According to the reviewer recommendations, previous figure 1 and 3 have been combined into one (new figure no. 1) and figure 2 has been combined with figure 4 to make new figure no. 2. The main text has also been amended accordingly.

**Comment:** F11 and F12: Please delete...instead add LC values tables
Response: The LC values have already been added in tables 1 and 2. The figures 11 and 12 provide the visual demonstration of the larvicidal activity, so with the kind consent of our proficient reviewer we would like to keep them in the main text.

References:

R1. Abu-Darwish MS, Efferth T. 2018 Medicinal plants from near east for cancer therapy. Front. Pharmacol. 9, 56. (doi: 10.3389/fphar.2018.00056)

R2. Kadir SLA, Yaakob H, Mohamed RZ. 2013 Potential anti-dengue medicinal plants: a review. J Nat Med 67:677–689 (doi: 10.1007/s11418-013-0767-y)

R3. Govindachari TR, Suresh G, Gopalakrishnan G, Banumathy B, Masilamani S. 1998 Identification of antifungal compounds from the seed oil of Azadirachta indica. Phytoparasitica 26, 109–116. (doi: 10.1007/BF02980677)

R4. Shawkey AM, Abdulall AK, Rabeh MA, Abdellatif AO. 2014 Enhanced biocidal activities of Citrullus colocynthis aqueous extracts by green nanotechnology. International Journal of Applied Research in Natural Products 7 (2), 1-10.

R5. Shawkey AM, Rabeh MA, Abdulall AK, Abdellatif AO. 2013 Green nanotechnology: Anticancer activity of silver nanoparticles using Citrullus colocynthis aqueous extracts. Adv. Life Sci. Technol. 13, 60–70.

R6. Rawani A, Ghosh A, Chandra G. 2013 Mosquito larvicidal and antimicrobial activity of synthesized nano-crystalline silver particles using leaves and green berry extract of Solanum nigrum L. (Solanaceae: Solanales). Acta Tropica 128, 613–622. (doi: 10.1016/j.actatropica.2013.09.00)

R7. Sujitha V, Murugan K, Paulpandi M, Panneerselvam C, Suresh U, Roni M, Nicoletti M, Higuchi A, Madhiyazhagan P, Subramaniam J, Dinesh D, Vadivalagan C, Chandramohan B, Alarfaj AA, Munusamy MA, Barnard DR, Benelli G. 2015 Green-synthesized silver nanoparticles as a novel control tool against dengue virus (DEN-2) and its primary vector Aedes aegypti. Parasitol Res., 114, 3315-25. (doi: 10.1007/s00436-015-4556-2)

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R9. Suresh U, Murugan K, Benelli G, Nicoletti M, Barnard DR, Panneerselvam C, Kumar PM, Subramaniam J, Dinesh D, Chandramohan B. 2015 Tackling the growing threat of dengue: Phyllanthus niruri-mediated synthesis of silver nanoparticles and their mosquitocidal properties against the dengue vector Aedes aegypti (Diptera: Culicidae). Parasitol Res. 114(4), 1551-1562. (doi: 10.1007/s00436-015-4339-9)
R10. Kanwal Z, Raza MA, Riaz S, Manzoor S, Tayyeb A, Sajid I, Naseem S. 2019 Synthesis and characterization of silver nanoparticle-decorated cobalt nanocomposites (Co@AgNPs) and their density-dependent antibacterial activity. *R. Soc. open sci.* 6, 182135. (doi:10.1098/rsos.182135)

R11. Kanwal Z, Raza, MA, Manzoor F, Riaz S, Jabeen G, Fatima S, Naseem S. 2019 A Comparative Assessment of Nanotoxicity Induced by Metal (Silver, Nickel) and Metal Oxide (Cobalt, Chromium) Nanoparticles in *Labeo rohita*. *Nanomaterials*, 9, 309. (doi:10.3390/nano9020309)

R12. 22-Aparajita V, Mohan SM. 2016 Controllable synthesis of silver nanoparticles using Neem leaves and their antimicrobial activity. *J Radiat Res Appl Sc* 9(1), 109-115. (doi: 10.1016/j.jrras.2015.11.001)

R13. 23-Shawkey AM, Rabeh MA, Abdulall AK, Abdellatif AO. 2013 Green nanotechnology: Anticancer activity of silver nanoparticles using Citrullus colocynthis aqueous extracts. *Adv. Life Sci. Technol.* 13, 60–70.

R14. Devadiga A, Vidya SK, Saidutta MB. 2017 Effect of Precursor Salt Solution Concentration on the Size of Silver Nanoparticles Synthesized Using Aqueous Leaf Extracts of *T. catappa* and *T. grandis* Linn f.—A Green Synthesis Route. In: Mohan B. R., Srinikethan G., Meikap B. (eds) Materials, Energy and Environment Engineering. Springer, Singapore. (doi: 10.1007/978-981-10-2675-1_17)

R15. Vishwakarma K, Shweta, Upadhyay N, Singh J, Liu S, Singh VP, Prasad SM, Chauhan DK, Tripathi DK, Sharma S. 2017 Differential Phytotoxic Impact of Plant Mediated Silver Nanoparticles (AgNPs) and Silver Nitrate (AgNO₃) on *Brassica sp*. *Front Plant Sci* 8, 1501. (doi: 10.3389/fpls.2017.01501)

R16. Subramaniam K, Siswomihardjo W, Sunarintyas S. 2005 The effect of different concentrations of Neem (Azadiractha indica) leaves extract on the inhibition of *Streptococcus mutans* (In vitro). *Maj Ked Gigi (Dent J)* 38, 176-9. ...

R17. Ederley V, Gloria C, Gladis M, César H, Jaime O, Oscar A. 2018 Silver Nanoparticles Obtained by Aqueous or Ethanolic *Aloe vera* Extracts: An Assessment of the Antibacterial Activity and Mercury Removal Capability. *J Nanomater* 2018, Article ID 7215210, 7 pages (doi: 10.1155/2018/7215210)
Upper panel-Different steps during the preparation of aqueous extract of *A. indica* leaves; Middle Panel-Synthesis of AgNPs; Lower Panel- (h) precursor solution, (i) reducing agent, (j-l) colloidal samples of AgNPs prepared by different amounts of *A. indica* extract.

229x252mm (150 x 150 DPI)
Upper panel-Different steps during the preparation of aqueous extract of C. colocynthis leaves; Middle Panel- Synthesis of AgNPs; Lower Panel- (h) precursor solution, (i) reducing agent, (j-l) colloidal samples of AgNPs prepared by different amounts of C. colocynthis extract.

257x248mm (150 x 150 DPI)
Schematic illustration of the synthesis mechanism of extract-mediated AgNPs. The phenomena occurring in each phase is mentioned accordingly.

286x153mm (150 x 150 DPI)
UV-Vis spectroscopy results of AgNPs synthesized by (a) three different concentrations of A. indica leaves extract and (b) three different concentrations of C. colocynthis fruit extract. Single absorption peak in each case appeared in the wavelength range of 412-430 nm indicating the spherical shaped AgNPs. Insets display images of each colloidal sample in cuvette.
SEM micrographs and histograms. (a,b) SEM image of A. indica AgNPs showed spherical shaped NPs decorated on larger extract particles, (c) Histogram of A. indica AgNPs indicated their diameter ranging between 12-24 nm. (d,e) SEM results of C. colocynthis AgNPs reveal the sphere like morphology and random dispersal over the bigger extract particles. (f) Histogram of C. colocynthis AgNPs indicated their diameter ranging between 20-36 nm.

321x187mm (150 x 150 DPI)
EDX spectrum showing elemental composition analysis of AgNPs (a) *A. indica* leaves, (b) *C. colocynthis* fruit. Insets show the table of elements in each case.

256x303mm (150 x 150 DPI)
X-ray diffraction patterns of (a) A. indica AgNPs and (b) C. colocynthis AgNPs
FTIR spectra (a) A. indica AgNPs, (b) C. colocynthis AgNPs. Occurrence of various absorption peaks at different positions indicates the presence of different biomolecules on the surface of NPs.
A representative display of larvicidal bioassay. Water (control), A. indica extract (Conc: 1.5 %), AgNPs at different concentrations. Upper panel shows the initial stage of experiment with 0 h time. Lower panel shows results after 24 h.

370x118mm (150 x 150 DPI)
A typical demonstration of larvicidal activity. Water (control), C. colocynthis extract (Conc: 1.5 %), AgNPs at different concentrations. Upper panel shows stage in the start of the experiment (at 0 h). Lower panel displays the final stage story (after 24 h).
Graphs showing the larvicidal effect of AgNPs at different concentrations (a) A. indica AgNPs showed LC50 at about 1.25 mg/L while at 20 mg/L 100% mortality was observed, (b) C. colocynthis AgNPs exhibited LC50 at 0.3 mg/L and 100% mortality was observed at 5 mg/L.
Appendix D

Reply to Reviewer’s Reports (Manuscript ID RSOS-200540)

We are again grateful to the reviewers for their skillful suggestions and valuable comments to improve our manuscript. All minors points and issues raised by the learnt reviewers have completely been addressed and the manuscript has been modified accordingly. A detailed point-by-point response to all comments is also provided below indicating the implemented changes in in the revised version of the manuscript. A highlighted version by ‘Track Changes’ is also included with the resubmission.

Response to Comments Reviewer # 2

Comment: I appreciate the efforts taken in revision of the manuscript. A few errors are yet to be addressed. The errors have been incorporated in the MS. Authors are requested to go through the MS and address them.

Response: We are obliged to the expert reviewer for appreciating our efforts in revised manuscript. All the issues and errors highlighted by the reviewer are addressed and the main text has been modified accordingly.

Comment: “only AgNPs prepared by using the 500 µL of extract were taken as optimized NPs” Can you provide a reason for choosing this extract concentration

Response: We chose 50 mL A. indica extract mediated AgNPs for bioassay on the basis of UV-Vis spectroscopy results. It can be seen from Uv-Vis spectra (Figure 4a) that the broadening of absorption peaks for 250 µL and 1000 µL extract mediated AgNPs were larger as compared to 500 µL AgNPs. Narrow peak width qualitatively indicates monodispersity and uniformity in the size distribution of NPs. This has also been mentioned in the main text.

Comment: Can you provide a reason for choosing this particular (2 mL extract mediated) NPs

Response: In the case of C. colocynthis extract mediated AgNPs, again on the basis of Uv-Vis spectra, 2 mL extract mediated AgNPs were selected for larvicidal activity. Because of strong absorption intensity and narrow peak width as compared to 1 mL and 4 mL extracted mediated AgNPs respectively. The main text has also been modified to rectify this point.