

**Dr. Vijai K. Rai**  
**Professor**



**Department of Chemistry**

**University of Lucknow**

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## Education:

Ph. D. (2006) Department of Chemistry, University of Allahabad, U. P. India.

UGC-CSIR-NET (Chemical Sciences) qualified.

M. Sc. (72.2 %) From DDU Gorakhpur University, U. P. India.

B. Sc. (71.9 %) From DDU Gorakhpur University, U. P. India.

## Experience:

- ❖ Assistant Professor at SMVD University, Jammu, w.e.f. 20. 08. 2009.
- ❖ Assistant Professor at Guru Ghasidas Vishwavidyalaya, Bilaspur, C.G. w.e.f. 09. 08. 2011.
- ❖ Postdoctoral Research Work (From 01. 04. 2007 to 19. 04. 2009).
- ❖ Worked at the **University of Calgary, CANADA**

## Awards Received

- |    |                                    |                             |  |
|----|------------------------------------|-----------------------------|--|
| 1. | Fast Track Young Scientist Award   | 4 <sup>th</sup> March 2011  | <b>DST, Government of India</b> , New Delhi, INDIA |
| 2. | Young Scientist Award              | 10 <sup>th</sup> Feb. 2010  | 5 <sup>th</sup> J K Science Congress, Jammu, India |
| 3. | <b>Golden Jubilee Award (NASI)</b> | 21 <sup>st</sup> Nv., 2008  | <b>National Academy of Sciences</b> , India        |
| 4. | D. S. Bhakuni Award                | 26 <sup>th</sup> Dec., 2007 | <b>Indian Chemical Society</b> , India             |
| 5. | Young Scientist Award              | 5 <sup>th</sup> Fe., 2007   | International Academy of Physical Sciences, India  |

## Completed Research Projects – 03 (50 Lakhs)

- |    |                 |   |
|----|-----------------|---|
| 1. | Funding Agency: | <b>University Grants Commission (UGC), New Delhi, India</b>                                   |
|    | Ref No.:        | F. No. 39-764/2010 (SR)   |
|    | Title:          | <i>Access to potentially antiviral novel nucleosides using microwave methodology</i>          |
| 2. | Funding Agency: | <b>Council of Scientific &amp; Industrial Research (CSIR), New Delhi, India</b>               |
|    | Ref No.:        | No. 01 (2442)/10/(EMR-II)   |
|    | Title:          | <i>Access to novel imino-/thiosugar scaffolds from renewable bioresources</i>                 |
| 3. | Funding Agency: | <b>Department of Science &amp; Technology (DST), New Delhi, India</b>                         |
|    | Ref No.:        | No. SR/FT/CS-99/2010  |
|    | Title:          | <i>NHC-/enamine-iminium catalysis in stereocontrolled construction of bioactive scaffolds</i> |

## Ongoing Research Project – 01 (46 Lakhs)

- Funding Agency: **SERB, DST, New Delhi, India**  
 Ref No.: **CRG/2021/001162**  
 Title: *Photoredox Catalysis to Access Stereoselective Cascade Reactions*  
 Start Date: 02 December 2021

## Research Interest

My research basically bioactive compounds, which are important pharmaceutical and drug candidates. However, most importantly *my work is based on environmental-friendly green protocols that reduce or eliminate hazardous chemicals and wastes and use bio-renewable resources as starting materials in some cases or using nano-materials and visible light as catalysts.*

- ❖ Heterogeneous Catalysis using nano-materials
- ❖ Visible-Light Induced Organic Reactions
- ❖ Stereo-controlled Construction of C-C and C-Hetero Bond
- ❖ Small & Medium Ring Heterocyclic Syntheses
- ❖ Homogeneous Catalysis, Ionic Liquids, Organocatalysis

## Ph D Supervision – Thesis Awarded

The following two (02) Ph D students have been awarded their Ph. D. degrees under my supervision:

- Suhasini Mahata**: Degree awarded on 05 August 2019.
- Fooleswar Verma**: Degree awarded on 21 August 2019.

## Ph D Supervision – Currently Working

Following two (02) Ph D students are working on their Ph. D. degrees and one (01) as JRF in SERB-CRG Project under my supervision: **1. Ambika Asati** **2. Bhushashi Khunte**

## Member of Academic/Professional Forum

- ❖ Editorial Board Member of *Letters in Organic Chemistry*, an International Journal
- ❖ Life Member of the Indian Science Congress.

## Book Chapters Published (02):

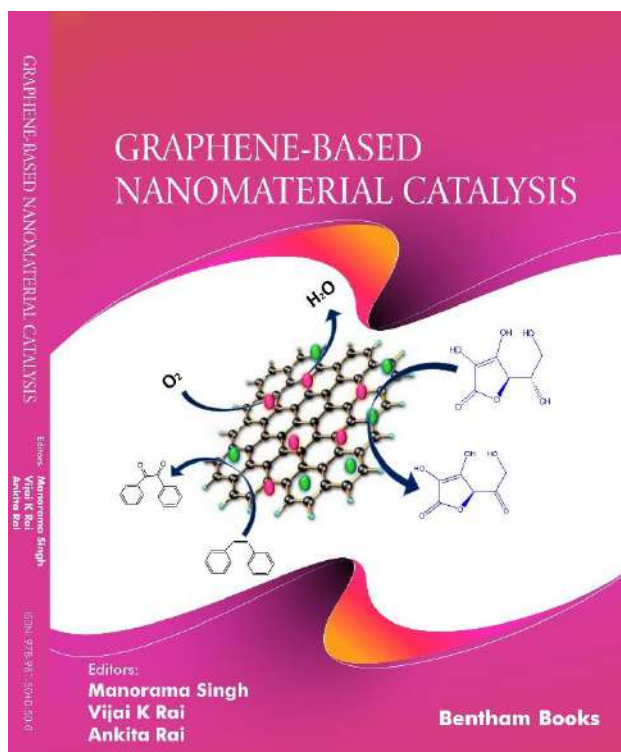
### 1. *Role of MOFs as Electro/-Organic Catalysts*

M. Singh, A. Rai, **Vijai K. Rai**, S. R. Bhardiya, A. Asati, Applications of Metal-Organic Frameworks and their derived materials, **2020**; ISBN 978-1-119-65098-0. (Wiley-Scrivener Publishing, Beverly, MA).

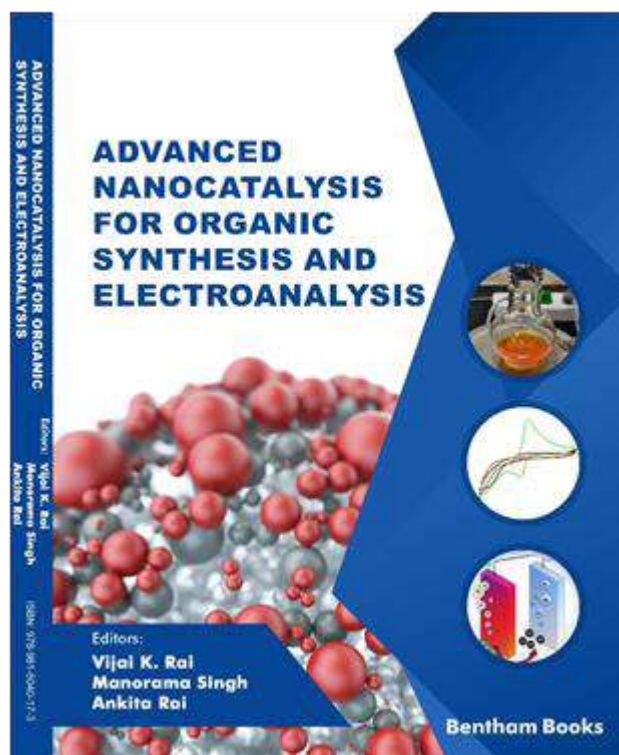
### 2. *Electrocatalysis: Application of nanocomposite materials*

M. Singh, A. Rai, **Vijai K. Rai**, Methods for Electrocatalysis: Advanced Materials and Allied Applications, **2020** ISBN 978-3-030-27161-9. (Springer Nature, Switzerland)

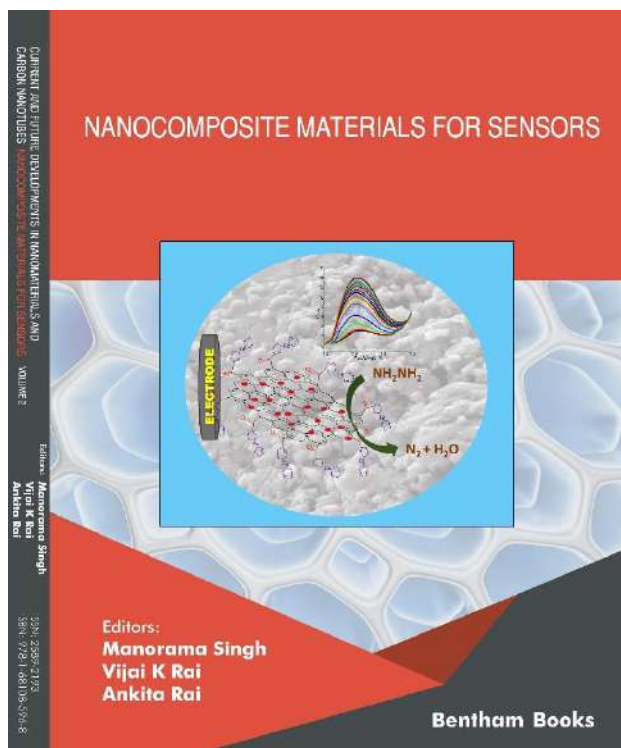
## Books Published (03):



1. *Graphene-Based Nanomaterial Catalysis*, 2022  
Bentham Science Publishers.  
ISBN (online): 978-981-5040-49-4



2. *Advanced Nanocatalysis for Organic Syntheses and Electroanalyses*, 2022, Bentham Science Publishers., ISBN (online): 978-981-5040-16-6



3. *Nanocomposite Materials for Sensors*, 2022  
Bentham Science Publishers.  
ISBN (online): 978-1-68108-596-8; ISSN  
(online): 2589-2193

## Research Publications

## Year 2022:

1. P. Shukla, M. Singh, **Vijai K. Rai**, A. Rai  
*Regioselective installation of enolizable ketones and unprotected mercaptoacetic acid into olefins using GO as phase transfer catalyst*  
*New J. Chem.* **2022**, 46, 329. **IF: 3.925**
2. M. Singh, S. R. Bhardiya, A. Rai, **Vijai K. Rai**  
*Electrochemical approach for recognition and quantification of p-phenylenediamine: a review*  
*Sens. Diagn.*, **2022**, DOI:10.1039/d1sd00070e.
3. M. Singh, S. R. Bhardiya, A. Rai, **Vijai K. Rai**  
*Graphene-based Nanomaterials for Electrochemical Sensing of Hydrazine: A Review*  
*Curr. Anal. Chem.*, **2022**, (In Press). **IF: 2.374**

## Year 2021:

4. *Design of a Sensitive Electrochemical Sensor Based on Ferrocene-Reduced Graphene Oxide/Mn-Spinel for Hydrazine Detection*  
M. Singh, S. R. Bhardiya, A. Asati, H. Sheshma, A. Rai, **Vijai K. Rai**  
*Electroanalysis*, **2021**, 33, 464-472. **IF: 3.077**
5. *Sensitive electrocatalytic determination of p-phenylenediamine using bimetallic nanocomposite of Cu-Ag nanoalloy and ionic liquid-graphene oxide*  
M. Singh, S. R. Bhardiya, A. Asati, H. Sheshma, **Vijai K. Rai**, A. Rai,  
*J Electroanal Chem.* **2021**, 894, 115360-115368. **IF: 4.598**
6. *A novel bioconjugated reduced graphene oxide-based nanocomposite for sensitive electrochemical detection of cadmium in water*  
S. R. Bhardiya, A. Asati, H. Sheshma, A. Rai, **Vijai K. Rai**, M. Singh  
*Sensors & Actuators: B. Chemical*, **2021**, 328, 129019-129028. **IF: 9.221**

## Year 2020:

7. *Facile Synthesis of  $\gamma$ -Ketonitriles in Water via C(sp<sup>2</sup>)-H Activation of Aromatic Aldehydes over Cu@g-C<sub>3</sub>N<sub>4</sub> under Visible-Light*  
**Vijai K. Rai**, F. Verma, S. R. Bhardiya, H. Sheshma, A. Rai, M. Singh  
*Eur. J. Org. Chem.*, **2020**, 5841-5846. **IF: 3.261**
8. *Metal-Free C-H Activation over Graphene Oxide Towards Direct Syntheses of Structurally Different Amines and Amides in Water*  
P. Shukla, A. Asati, S. R. Bhardiya, M. Singh, **Vijai K. Rai**, A. Rai  
*J. Org. Chem.*, **2020**, 85, 15551-15561. **IF: 4.198**
9. *Cu/Cu<sub>2</sub>O@g-C<sub>3</sub>N<sub>4</sub>: Recyclable Photocatalyst under Visible-Light to Access 2-Aryl-/benzimidazoles/benzothiazoles in Water*  
P. K. Singh, S. R. Bhardiya, A. Asati, **Vijai K. Rai**, M. Singh, A. Rai  
*Chemistry Select*, **2020**, 5, 14270-14275. **IF: 2.307**
10. *Cu (I)-Induced Activation of Furan for Inverse Electron Demand ADAR with Alkenes towards Regioselective Synthesis of Tetrahydropyridine*, P. Shukla, A. Asati, S. R. Bhardiya, M. Singh, **Vijai K. Rai**, A. Rai  
*J. Org. Chem.*, **2020**, 85, 7772-7780. **IF: 4.198**

11. *A Novel Ternary Graphene-Based Nanocomposite Modified Electrode for Acetaminophen Detection*  
M. Singh, A. Sahu, P. K. Singh, F. Verma, A. Rai, **Vijai K. Rai**  
*Electroanalysis*, **2020**, 32, 1516-1522. **IF: 3.077**
12. *Bio-reduction of Graphene Oxide: catalytic applications of (reduced) GO in organic synthesis*  
**Vijai K. Rai**, S. Mahata, H. Kashyap, M. Singh, A. Rai  
*Curr. Org. Synth.*, **2020**, 17, 164-191. **IF: 2.276**
13. *Graphene-based Nanomaterials for Fabrication of 'Pesticide' Electrochemical Sensors*  
M. Singh, S. R. Bhardiya, F. Verma, **Vijai K. Rai**, A. Rai  
*Current Graphene Science*, **2020**, 3, 26-40.

## Year 2019:

14. *Photocatalytic C(sp<sup>3</sup>)-H activation towards  $\alpha$ -methylenation of ketones using MeOH as IC source steering reagent*  
F. Verma, P. Shukla, S. R. Bhardiya, M. Singh, A. Rai, **Vijai K. Rai**  
*Adv. Synth. Catal.* **2019**, 361, 1247-1252. **IF: 5.981**
15. *Metal Doped-C<sub>3</sub>N<sub>4</sub>/Fe<sub>2</sub>O<sub>4</sub>: Efficient and Versatile Heterogenous Catalysts for Organic Transformations*  
**Vijai K. Rai**, F. Verma, S. Mahata, S. R. Bhardiya, M. Singh, A. Rai  
*Curr. Org. Chem.*, **2019**, 23, 1282-1304. **IF: 2.226**
16. *Efficient electrochemical determination of p-aminophenol using a novel tricomponent graphene-based nanocomposite*  
M. Singh, A. Sahu, S. Mahata, P. K. Singh, A. Rai, **Vijai K. Rai**  
*New J. Chem.*, **2019**, 43, 14972-14978. **IF: 3.925**
17. *A facile and efficient carbocatalytic route to quaternary C-bearing N-tosylaziridines from Morita-Baylis-Hillman adduct in water*  
P. Shukla, S. Mahata, H. Kashyap, M. Singh, **Vijai K. Rai**, A. Rai  
*Tetrahedron Lett.* **2019**, 60, 1943-1948. **IF: 2.032**
18. *Efficient electrocatalytic oxidation of p-phenylenediamine using a novel PANI/ZnO anchored bio-reduced graphene oxide nanocomposite*  
M. Singh, A. Sahu, S. Mahata, P. Shukla, A. Rai, **Vijai K. Rai**  
*New J. Chem.* **2019**, 43, 6500-6506. **IF: 3.925**
19. *A novel carbocatalytic hydride transfer strategy for efficient reduction of structurally different aldehydes and ketones in water*  
**Vijai K. Rai**, S. Mahata, S. R. Bhardiya, P. Shukla, A. Rai, M. Singh  
*Tetrahedron Lett.*, **2019**, 60, 524-529. **IF: 2.032**
20. *AuNPs/Neutral red-biofunctionalized graphene nanocomposite for nonenzymatic electrochemical detection of organophosphate via NO<sub>2</sub> reduction*  
M. Singh, H. Kashyap, P. K. Singh, S. Mahata, **Vijai K. Rai**, A. Rai  
*Sensors & Actuators: B. Chemical*, **2019**, 290, 195-202. **IF: 9.221**
21. *Visible Light-Induced Direct Conversion of Aldehydes into Nitriles in Aqueous Medium Using Co@g-C<sub>3</sub>N<sub>4</sub> as Photocatalyst*  
F. Verma, P. Shukla, S. R. Bhardiya, M. Singh, A. Rai, **Vijai K. Rai**  
*Catalysis Commun.* **2019**, 119, 76-81. **IF: 3.510**



22. *A Facile Iodine-Promoted Enal-Based cis-Selective Construction of Aziridin-2-aldehyde in Water*  
P. K. Singh, F. Verma, S. R. Bhardiya, M. Singh, **Vijai K. Rai**, A. Rai  
*Chemistry Select*, **2019**, 4, 1240-1243. **IF: 2.307**
23. *First bio-covalent functionalization of graphene with threonine towards drug sensing via electrocatalytic transfer hydrogenation*  
A. Sahu, P. Shukla, S. Mahata, **Vijai K. Rai**, A. Rai, M. Singh  
*Sensors & Actuators: B. Chemical*, **2019**, 281, 1045-1053. **IF: 9.221**

## Year 2018:

24. *A novel and efficient reduction of graphene oxide using Ocimum sanctum L. leaf extract as an alternative renewable bio-resource*  
S. Mahata, A. Sahu, P. Shukla, A. Rai, M. Singh, **Vijai K. Rai**  
*New J. Chem.* **2018**, 42, 19945-19952. **IF: 3.925**
25. *Visible-light driven regioselective synthesis of 1H-tetrazoles from aldehydes through isocyanide-based [3+2] cycloaddition*  
F. Verma, A. Sahu, P. K. Singh, A. Rai, M. Singh, **Vijai K. Rai**  
*Green Chem.* **2018**, 20, 3783-3789. **IF: 11.034**
26. *One-Pot Allan–Robinson/Friedländer Route to Chromen-/Quinolin-4-ones through the Domino Acetylation Cyclisation of 2-Hydroxy-/2-Aminobenzaldehyde*  
**Vijai K. Rai**, F. Verma, G. P. Sahu, M. Singh, A. Rai  
*Eur. J. Org. Chem.* **2018**, 537–544. **IF: 3.261**
27. *Morita-Baylis-Hillman enal-based triple cascade strategy for anti-selective synthesis of highly functionalized tetrahydropyridines using iminium-enamine catalysis*  
**Vijai K. Rai**, F. Verma, M. Satnami, M. Singh, A. Rai  
*Tetrahedron Lett.*, **2018**, 59, 1783–1786. **IF: 2.023**
28. *Aqueous mortar-pestle grinding: An efficient, attractive, and viable technique for the regioselective synthesis of  $\beta$ -amino alcohols*  
N. Singh, **Vijai K. Rai**, A. Kumar  
*Competes Rendus Chimie*, **2018**, 21, 71-79. **IF: 2.550**
29. *Bio-inspired unprecedented synthesis of reduced graphene oxide: a catalytic probe for electro-/chemical reduction of nitro groups in an aqueous medium*  
S. Mahata, A. Sahu, P. Shukla, A. Rai, M. Singh, **Vijai K. Rai**  
*New J. Chem.*, **2018**, 42, 2067-2073. **IF: 3.925**
30. *Graphene oxide catalyzed C-N/C-S/[3+2] cyclization cascade for green synthesis of thiazolidinone in water*  
S. Mahata, A. Sahu, P. Shukla, A. Rai, M. Singh, **Vijai K. Rai**  
*Lett. Org. Chem.* **2018**, 15, 665-672. **IF: 0.797**

## Year 2017:

31. *A co-operative effect of visible light photo-catalysis and CoFe<sub>2</sub>O<sub>4</sub> nanoparticles for green synthesis of furans in water*  
F. Verma, P. K. Singh, S. R. Bhardiya, M. Singh, A. Rai, **Vijai K. Rai**  
*New J. Chem.* **2017**, 41, 4937-4942. **IF: 3.925**

32. *Facile construction of AuNPs modulated SDS wrapped G-TC tailored electrode for sensitive detection of ascorbic acid*  
H. Kashyap, P. K. Singh, F. Verma, **Vijai K. Rai**, A. Rai, M. Singh  
*New J. Chem.* **2017**, 41, 6489-6496. **IF: 3.925**
33. *First graphene oxide promoted metal-free nitrene insertion into olefins in water: towards facile synthesis of activated aziridines*  
P. Shukla, S. Mahata, A. Sahu, M. Singh, **Vijai K. Rai**, A. Rai  
*RSC Advances*, **2017**, 7, 48723-48729. **IF: 4.036**
34. *Synthesis of 2-amino-4H-chromen-4-ylphosphonates and  $\beta$ -phosphonomalonates via tandem Knoevenagel-Phospha-Michael reaction and antimicrobial evaluation of newly synthesized  $\beta$ -phosphonomalonates.*  
P. Kour, A. Kumar, R. Sharma, R. Chib, I. A. Khan, **Vijai K. Rai**  
*Res Chem. Intermed.* **2017**, 43, 7319-7329. **IF: 3.134**
35.  *$\text{Bi}(\text{NO}_3)_3 \cdot 5\text{H}_2\text{O}$  catalyzed phosphorylation of aldehydes: an efficient route to  $\alpha$ -hydroxyphosphonates*  
A. Kumar, S. Jamwal, S. Khan, N. Singh, **Vijai K. Rai**  
*Phosphorus, Sulfur, Silicon and The Related Elements* **2017**, 192, 381-385. **IF: 1.051**
36. *Aqueous microwave-assisted DMAP catalyzed synthesis of  $\beta$ -phosphonomalonates and 2-amino-4H-chromen-4-ylphosphonates via a domino Knoevenagel-phospha-Michael reaction*  
P. Kour, A. Kumar, **Vijai K. Rai**  
*Competes Rendus Chimie*, **2017**, 20, 140-145. **IF: 2.550**
37. *One-pot synthesis of highly functionalized pyrido-1,3-thiazin-4-ones using unprotected sugars in a Task-specific Ionic Liquid, [Bmim]SCN*  
**Vijai K. Rai** & V. R. Sharrof  
*J. Heterocyclic Chem.*, **2017**, 54, 1178-1185. **IF: 2.035**

## Year 2016:

38. *First iodine/IL-catalyzed carbohydrate activation as aldehyde equivalent for  $[\text{C}+2\text{C}+\text{N}]$  construction of  $\beta$ -lactam ring*  
**Vijai K. Rai**, B. Sharma, V. R. Sharoff, A. Rai,  
*Tetrahedron Lett.* **2016**, 57, 3260-3263. **IF: 2.032**
39. *One-pot cis-selective route to sugar-fused thiazines via a masking-unmasking strategy in basic ionic liquid*  
**Vijai K. Rai**, Rahul K. Kosta,  
*Can. J. Chem.* **2016**, 94, 827-832. **IF: 1.051**
40. *Carbocation catalyzed carboxylic acid activation in Staudinger reaction for stereoselective synthesis of  $\beta$ -lactams,*  
A. Rai, P. K. Singh, P. Shukla, **Vijai K. Rai**,  
*Tetrahedron Lett.* **2016**, 57, 5084-5088. **IF: 2.032**
41. *Decoration of GO with Fe spinel-Naf/DMAP: an electrochemical probe for sensing  $\text{H}_2\text{O}_2$  reduction,*  
M. Singh, S. R. Bhardiya, H. Kashyap, F. Verma, **Vijai K. Rai** & I. Tiwari,  
*RSC Advances*, **2016**, 6, 104868-104874. **IF: 4.036**
42. *A facile anti-selective synthesis of 3-nitropyridin-2-ones using Morita-Baylis Hillman adduct of nitroalkene*  
**Vijai K. Rai**, G. P. Sahu, M. Singh & A. Rai  
*Lett. Org. Chem.*, **2016**, 13, 547-553. **IF: 0.797**

## Year 2015-14:

43. *The first NHC-induced regioselective introduction of C- and N-nucleophiles in to Baylis–Hillman enals*,  
Vijai K. Rai, G. P. Sahu, A. Rai  
*Tetrahedron Lett.* **2015**, 55, 2664-2668. **IF: 2.032**

## Year 2013:

44. *Masked amino acid: a new C-nucleophile for I<sub>2</sub>-catalyzed stereoselective ring opening of epoxides in ionic liquid*  
Vijai K. Rai, R. Sharma, A. Kumar  
*Tetrahedron Lett.* **2013**, 54, 1071-1075. **IF: 2.032**
45. *The First I<sub>2</sub> Promoted Efficient Aminoacetylation of Activated Aziridines in Ionic liquids*,  
Vijai K. Rai, N. Sharma & A. Kumar,  
*Synlett* **2013**, 24, 097-101. **IF: 2.170**
46. *CeCl<sub>3</sub>·7H<sub>2</sub>O/NaI-Promoted Direct Synthesis of 1,3-Benzoxazine-2-Thione N-Nucleosides under Microwave Irradiation*  
Vijai K. Rai, N. Singh  
*Nucleosides, Nucleotides & Nucleic Acids* **2013**, 32, 247-255. **IF: 1.449**
47. *Masked mercapto acid-driven MCR in task-specific ionic liquid: a new stereocontrolled entry into bicyclic 1,3-thiazines*  
Vijai K. Rai, P. K. Rai, Y. Thakur  
*Tetrahedron Lett.* **2013**, 54, 6469-6473. **IF: 2.032**

## Year 2012:

48. *Efficient Ce(III)-Catalyzed Cis-Selective Synthetic Approach to  $\gamma$ -Lactones in Aqueous Media*,  
Vijai K. Rai, P. Tikku, A. Kumar;  
*Synth. Commun.* **2012**, 42, 1489-1499. **IF: 1.937**

## Year 2011:

49. *An unprecedented synthesis of  $\gamma$ -lactams via mercaptoacetylation of aziridines in water*,  
Vijai K. Rai, P.K. Rai, S. Bajaj, A. Kumar  
*Green Chem.* **2011**, 13, 1217-1223. **IF: 11.034**
50. *[2 + 2] Annulation of aldimines with sulfonic acids: a novel one-pot cis-selective route to  $\beta$ -sultams*  
A. Rai, Vijai K. Rai, A. Singh, L.D.S. Yadav  
*Eur. J. Org. Chem.* **2011**, 4302-4306. **IF: 3.261**
51. *Nucleophilic acylation of  $\alpha$ -haloketones with aldehydes: an umpolung strategy for the synthesis of 1,3-diketones*  
S. Singh, P. Singh, Vijai K. Rai, R. Kapoor, L.D.S. Yadav  
*Tetrahedron Lett.* **2011**, 52, 125-128. **IF: 2.032**

## Year 2010:

52. *Carbohydrate Building Block in Ugi 3-Component Coupling Reaction: Convenient Annulation of Iminosugars on Imidazoles*,  
Vijai K. Rai, S. Singh, P. Singh, L.D.S. Yadav,  
*Synthesis* **2010** 4051-4056. **IF: 2.969**



53. *N*-Heterocyclic Carbene Catalyzed Cross-Coupling of Aromatic Aldehydes with Baylis-Hillman Bromides: An Easy Access to  $\alpha$ -Arylidene- $\gamma$ -keto Esters,  
P. Singh, S. Singh, **Vijai K. Rai**, L.D.S. Yadav,  
*Synlett* **2010**, 2649-2653. IF: 2.170
54. *Expedition synthesis of functionalized piperidines via NHC-catalyzed regioselective aziridine ring-opening with enals*, S. Singh, **Vijai K. Rai**, P. Singh, L.D.S. Yadav,  
*Synthesis* **2010**, 2957-2964. IF: 2.969
55. *NHC-catalyzed efficient synthesis of  $\beta'$ -amino enones via carbonyl umpolung reaction of enals with aziridines*, L.D.S. Yadav, **Vijai K. Rai**, S. Singh, P. Singh,  
*Tetrahedron Lett.* **2010**, 51, 1657-1662. IF: 2.032
56. *Carbonyl umpolung reactivity of enals: NHC-catalyzed synthesis of Aldol products via epoxide ring-opening*, L.D.S. Yadav, S. Singh, **Vijai K. Rai**, P. Singh;  
*Synlett* **2010**, 240-246. IF: 2.170

### Year 2009:

57. *A one pot [Bmim]OH mediated synthesis of 3-benzamidocoumarins*  
L.D.S. Yadav, S. Singh & **Vijai K. Rai**;  
*Tetrahedron Lett.* **2009**, 50, 2208-2212. IF: 2.032
58. *Novel catalyst-free, step and pot economic efficient mercaptoacetylative cyclisation in H<sub>2</sub>O: synthesis of 3-mercaptocoumarins*;  
L.D.S. Yadav, S. Singh & **Vijai K. Rai**;  
*Green Chem.* **2009**, 11, 878-882. IF: 11.034
59. *The First Cu(OTf)<sub>2</sub>-catalyzed synthesis of structurally novel bicyclic 1,3-oxazines via condensation-dehydrazinative ring transformation cascades*;  
L.D.S. Yadav, A. Rai, **Vijai K. Rai** & C. Awasthi;  
*J. Chem. Res.* **2009**, 520-526. IF: 1.097
60. *A novel one-pot stereoselective synthesis of N-protected  $\alpha$ -amino acids from Morita-Baylis-Hillman acetates*;  
L.D.S. Yadav, **Vijai K. Rai** & S. Singh;  
*Synlett* **2009**, 1423-1428. IF: 2.170
61. *The first regio- and diastereoselective direct introduction of  $\alpha$ -mercaptoacetic acid/amide units into Morita-Baylis-Hillman acetates*  
L.D.S. Yadav & **Vijai K. Rai**;  
*Tetrahedron Lett.* **2009**, 50, 2414-2419. IF: 2.032
62. *The first ionic liquid-promoted one-pot diastereoselective synthesis of 2,5-diamino-/2-amino-5-mercapto-1,3-thiazin-4-ones using masked amino/mercapto acids*;  
L.D.S. Yadav, **Vijai K. Rai** & B. S. Yadav;  
*Tetrahedron* **2009**, 65, 1306-1315. IF: 2.388

### Year 2008:

63. *A convenient CeCl<sub>3</sub>.7H<sub>2</sub>O/NaI-promoted synthesis of structurally novel and strained tricyclic  $\beta$ -lactams from hydrazines*,  
L.D.S. Yadav, **Vijai K. Rai**;  
*Tetrahedron Lett.* **2008**, 49, 5553-5556. IF: 2.032

64. *K-10 clay-catalyzed enol-driven decarboxylative ring transformation approach to dihydro- and tetrahydroquinolines from carbohydrates*,  
L.D.S. Yadav, C. Awasthi, **Vijai K. Rai**, A. Rai;  
*Synlett* **2008**, 2257-2262. IF: 2.170
65. *Diversity oriented synthesis of fused-ring 1,3-oxazines from carbohydrates as biorenewable feedstocks*  
L.D.S. Yadav, V.P. Srivastava, **Vijai K. Rai**, R. Patel;  
*Tetrahedron* **2008**, 64, 4246-4253. IF: 2.388
66. *A route to functionalized pyrimidines from carbohydrates via amine-driven dehydrative ring transformations*,  
L.D.S. Yadav, C. Awasthi, **Vijai K. Rai**, A. Rai;  
*Tetrahedron Lett.* **2008**, 49, 2377-2380. IF: 2.032
67. *Multicomponent reactions in chiral ionic liquids: a stereocontrolled route to mercaptopyranothiazoles*,  
L.D.S. Yadav, B.S. Yadav, **Vijai K. Rai**,  
*J. Heterocyclic Chem.* **2008**, 45, 1315-1319. IF: 2.035
68. *Carbohydrates to functionalized pyridines: a new synthetic approach via enol-driven ring transformations*,  
L.D.S. Yadav, A. Rai, **Vijai K. Rai**, C. Awasthi;  
*Synlett* **2008**, 529-534. IF: 2.170
69. *An expeditious synthesis of benzoxazine-2-thione C-nucleosides via Cu(OTf)<sub>2</sub>-mediated dehydrazinative  $\beta$ -glycosylation*;  
L.D.S. Yadav, **Vijai K. Rai**;  
*Nucleosides, Nucleotides & Nucleic Acids* **2008**, 27, 1227-1237. IF: 1.449
70. *Novel aziridination of  $\alpha$ -halo ketones: an efficient nucleophile-induced cyclization of phosphoramidates to functionalized aziridines*;  
L.D.S. Yadav, A. Rai, **Vijai K. Rai**, C. Awasthi;  
*Tetrahedron Lett.* **2008**, 49, 687-690. IF: 2.032
71. *Chiral ionic liquid-catalyzed Biginelli reaction: stereoselective synthesis of polyfunctionalized perhydropyrimidines*;  
L.D.S. Yadav, A. Rai, **Vijai K. Rai**, C. Awasthi;  
*Tetrahedron* **2008**, 64, 1420-1429. IF: 2.388

## Year 2007:

72. *A convenient synthesis of 1,2,4-trisubstituted azetidines by reductive cyclization of aza-Michael adducts of chalcones*.  
L.D.S. Yadav, C. Awasthi, **Vijai K. Rai**, A. Rai;  
*Tetrahedron Lett.* **2007**, 48, 8037-8039. IF: 2.032
73. *Multicomponent solvent-free cyclocondensation/glycosylation strategy for thiazolo-s-triazine N-nucleosides*;  
L.D.S. Yadav, **Vijai K. Rai**, S. Yadav,  
*Lett. Org. Chem.* **2007**, 4, 47-50. IF: 0.797
74. *Thiourea to dithiazolopyrimidines: highly regio- and stereoselective synthetic routes via mercaptoacetylation*;  
L.D.S. Yadav, S. Yadav, **Vijai K. Rai**;  
*Synthesis* **2007**, 3831-3838. IF: 2.969
75. *An efficient conjugate hydrothiocyanation of chalcones with a task-specific ionic liquid*;  
L.D.S. Yadav, R. Patel, **Vijai K. Rai**, V.P. Srivastava;  
*Tetrahedron Lett.* **2007**, 48, 7793-7795. IF: 2.032

76. *Biorenewable resources in the Biginelli reaction: Ce(III)-catalyzed synthesis of novel iminosugar-annulated perhydropyridines*;  
L.D.S. Yadav, A. Rai, **Vijai K. Rai**, C. Awasthi;  
*Synlett* **2007**, 1905-1908. **IF: 2.170**
77. *Biorenewable and mercaptoacetylating building blocks in the Biginelli reaction: synthesis of thiosugar-annulated dihydropyrimidines*;  
L.D.S. Yadav, C. Awasthi, **Vijai K. Rai**, A. Rai;  
*Tetrahedron Lett.* **2007**, 48, 4899-4902. **IF: 2.032**
78. *Thiourea to bicyclic scaffolds: highly regio/- and stereoselective routes to dithiazolopyrimidines*;  
L.D.S. Yadav, **Vijai K. Rai**;  
*Tetrahedron* **2007**, 63, 6924-6931. **IF: 2.388**
79. *One-pot dehydrazinative  $\beta$ -glycosylation in aqueous media: synthesis of benzoxazine C-nucleosides*;  
L.D.S. Yadav & **Vijai K. Rai**,  
*Synlett* **2007**, 1227-1230. **IF: 2.170**

## Year 2006:

80. *Chemoselective annulation of 1,3-dithiin/thiazine/oxathiin rings on thiazoles using green protocol*;  
L.D.S. Yadav & **Vijai K. Rai**,  
*Tetrahedron* **2006**, 62, 8029-8034. **IF: 2.388**
81. *A green protocol for annulation of s-triazine ring on thiazoles using three-component coupling strategy*;  
L.D.S. Yadav, S. Yadav & **Vijai K. Rai**,  
*Green Chem.* **2006**, 8, 455-458. **IF: 11.034**
82. *Novel mercaptoacetylative expeditious annulation of 5-mercaptopyrimidine ring on azoles using 1,3-oxathiolan-5-one*;  
L.D.S. Yadav, **Vijai K. Rai** & S. Yadav,  
*Tetrahedron* **2006**, 62, 5464-5468. **IF: 2.388**
83. *Active copper-promoted expeditious N-arylations in aqueous media under microwave irradiation*,  
L.D.S. Yadav, B. S. Yadav, **Vijai K. Rai**,  
*Synthesis* **2006**, 1868-1872. **IF: 2.969**
84. *Three-component coupling strategy for the expeditious synthesis of novel 4-aminobenzoxazinone N- nucleoside*;  
L.D.S. Yadav, **Vijai K. Rai**,  
*Tetrahedron Lett.* **2006**, 47, 395-397. **IF: 2.388**

## Year 2005:

85. *Mercaptoacetic acid based expeditious synthesis of polyfunctionalized 1,3-thiazines*  
L.D.S. Yadav & S. Yadav & **Vijai K. Rai**,  
*Tetrahedron* **2005**, 61, 10013-10017. **IF: 2.388**

## Year 2004:

86. *A novel salicylaldehyde-based mineral supported expedient synthesis of benzoxazinone nucleosides*,  
L.D.S. Yadav & B.S. Yadav, **Vijai K. Rai**,  
*Tetrahedron Lett.* **2004**, 45, 5351-5353. **IF: 2.032**

## Invited Lectures in Conferences

1. **Vijai K. Rai**, Visible-Light Induced Photoredox Catalysis in Cascade Reactions, 27th CONIAPS, October 26-28, 2021, (online mode) on Recent Advances in Chemical Sciences, jointly organized by SPS JNU and Dept of Chemistry BHU.
2. **Vijai K. Rai**, Design and Development of Visible-Light Photocatalytic Nanomaterials, Online Conference on Advanced Nano Materials (ICAN 2021), December 14-16, **2021**, Kerala.
3. **Vijai K. Rai**, Carbon Nitrides Under Visible-Light Conditions: Efficient Catalytic System for Organic Transformations, 26<sup>th</sup> CONIAPS, **18-20 December, 2020** at M. G. University, Kerala.
4. **Vijai K. Rai**, Visible Light Catalyzed- $\alpha$ -Methylenation of Ketones; 106<sup>th</sup> Indian Science Congress, **03-07 Jan., 2019** at Lovely Professional University, Punjab.
5. **Vijai K. Rai** Green Chemistry in Laboratory, Innovation in Engineering, C. G., **27-28 Feb. 2015** at J. K. Institute of Engineering, Bilaspur.
6. **Vijai K. Rai**, Green Chemistry: Why & How in Laboratory, In National Seminar on Chemistry in Our Lives, held on **1-2 Feb. 2013** at Science College, Bilaspur, Chhattisgarh
7. **Vijai K. Rai**, Green Synthetic Approaches in Drug Discovery Process, in Indian Chemical Society Conference held on **Dec. 12-15, 2012** at NITTTR, Shamla Hills, Bhopal, M. P.
8. **Vijai K. Rai**, CoFe<sub>2</sub>O<sub>4</sub>: Efficient Nano-catalyst for Facile Construction of Biomolecules & Role of H<sub>2</sub>O Towards Green Chemistry, In International Conference on Study of Nanomaterials and Scientific Development in 21<sup>st</sup> Century (ICSNSDC), **3-5 November, 2017** at Jiwaji University Gwalior.
9. **Vijai K. Rai**, Stringent and growing environmental regulation in Green Chemistry, In 12<sup>th</sup> CONIAPS, on **22-25 Dec, 2010** at University of Rajasthan, Jaipur.

## Paper Presented in Conferences

10. **Vijai K. Rai**, Metal Doped Carbon Nitrides: Efficient Heterogenous Catalysts for Organic Transformations, 26<sup>th</sup> ISCB, **22-24 January, 2020** at Nirma University, Ahmedabad.
11. **Vijai K. Rai**, Visible Light Induced [3+2] Cycloaddition: An Efficient Green Protocol for Tetrazoles; In International conference in Chemical Sciences including Green Chemistry; **27-29<sup>th</sup> December, 2018** at SRM Institute of Science & Technology, Ramapuram, Chennai.
12. **Vijai K. Rai**, Dual catalytic-effect of visible light and magnetic attraction for efficient synthesis of furan; In International conference on Global Trends in Pure and Applied Chemical Sciences; **8-9<sup>th</sup> December, 2017** at SRM University, Ghaziabad (U. P.).
13. **Vijai K. Rai**, Organocatalytic C-C and C-N Bond Forming Reactions Using Baylis-Hillman Adduct; oral presentation in 103<sup>rd</sup> Indian Science Congress, **03-07 January, 2016** at University of Mysore, Mysore.
14. **Vijai K. Rai**, Ion Implantation: A Green Chemistry Perspective; In National workshop, **18-19 Feb, 2014** at Guru Ghasidas Vishwavidyalaya, Bilaspur, C. G.
15. **Vijai K. Rai**, N-Heterocyclic Carbene-Catalyzed Efficient Synthesis of Functionalized Enones, In 5<sup>th</sup> JK Science Congress, on **8-10 Feb, 2010** at University of Jammu, J&K.

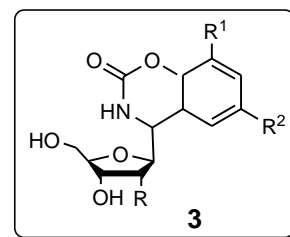
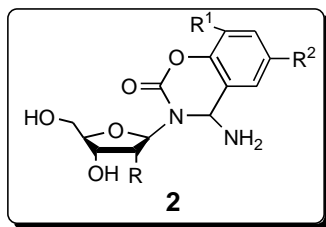
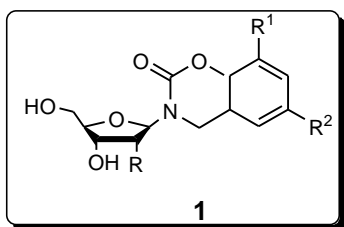
16. **Vijai K. Rai**; A Convenient  $\beta$ -Glycosylation of 1,3-Benzoxazine-2-thiones, organized by National Academy of Sciences (NASI), India, on **21-23 November, 2008** at Punjab University Chandigarh.
17. **Vijai K. Rai**; Chiral ionic liquid-catalyzed Biginelli reaction: stereoselective synthesis of polyfunctionalized perhydropyrimidines, presented in Proc. of 44<sup>th</sup> annual convention of chemist on **23-27 Dec. 2007** at Jaipur.
18. **Vijai K. Rai**; Chemoselective annulation of 1,3,4-thiadi- and dithiazine rings on imidazoles, presented in the 9<sup>th</sup> session of CONIAPS, **3-5 Feb. 2007** at Agra, India.
19. **Vijai K. Rai**; Green synthetic approach to thiazolo-s-triazine *N*-nucleosides, presented in the 76<sup>th</sup> annual session of NASI, **6-8 Oct. 2006** at IIT Bombay, India.
20. **Vijai K. Rai**; Multicomponent annulation of *s*-triazine ring on thiazoles under solvent-free microwave irradiation, In 93<sup>rd</sup> session of the Indian Science Congress, **3-7 Jan, 2006** at Hyderabad, India.

## Research-Summary

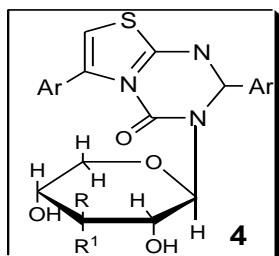
In my *over fourteen years of research experience in Synthetic Organic Chemistry*, I have been working mainly on synthesis and application in the following areas:

### [A]. Synthesis of Benzoxazinone C- and N-nucleosides:

Efavirenz (Sustiva), a benzoxazinone derivative, is a non-nucleoside reverse transcriptase inhibitor that has been approved by the FDA (September 17, 1998) and is presently in clinical use for the treatment of AIDS. In this respect, I have synthesized benzoxazinone *N*-nucleosides **1** (*Tetrahedron Lett.* **2004**, 45, 5351) and **2** (*Tetrahedron Lett.* **2006**, 47, 395) and their *C*-nucleosides **3** (*Synlett* **2007**, 1227). Compounds **1** were prepared by K-10 clay catalyzed cycloisomerization of salicylaldehyde 4-( $\beta$ -D-ribo- or  $\beta$ -D-2'-deoxyribofuranosyl) semicarbazones followed by reductive dehydrazination of their 4-hydrazinoderivatives. Compounds **2** were synthesized by K-10 clay supported three- component coupling reactions of substituted salicylaldehydes, ribosyl/deoxyribosylureas and ammonium acetate via cycloisomerisation of a aldimine intermediate. For benzoxazinone *C*-nucleosides **3**, a novel one-pot expeditious synthetic protocol has been developed via dehydrazinative  $\beta$ -glycosylation in aqueous media from the unprotected sugar and a compound containing an activated methylene group (Ref. 3). Recently, we have also reported 1,3-benzoxazine-2-thione *C*-nucleosides (*Nucleosides, Nucleotides & Nucleic Acids* **2008**, 27, 1227-1237).



### [B]. Synthesis of Glycon Modified N-nucleosides:

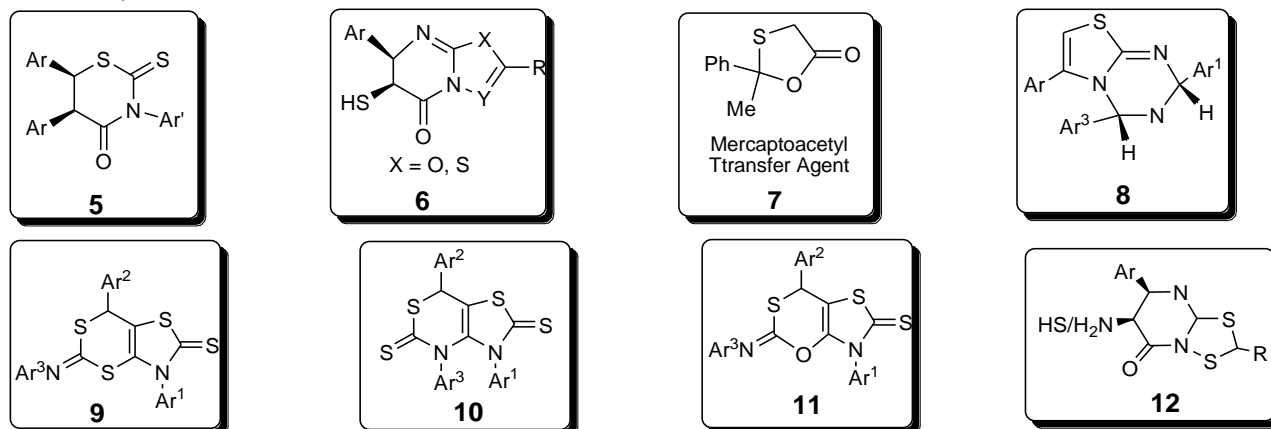


A green protocol involving novel three-component one-pot cyclocondensation reaction of 2-amino-4-aryl-thiazololes, aromatic aldehydes and ammonium thiocyanate under solvent-free MW irradiation conditions expeditiously yields thiazolo-s-triazine nucleobases, which afford the corresponding pyrano *N*-nucleosides **4** on iodine promoted glycosylation with 1,2,3,4-tetra-O-acetyl-beta-D-ribo-/xylopyranose under MW irradiation followed by deacetylation (Ref. 5).

**Ref. 5.** *Lett. Org. Chem.* **2007**, 4, 47.

### [C]. Stereo/-Chemo/- and Regioselective Syntheses:

Nowadays, selectivity in organic reactions, *i.e.* stereoselectivity, regioselectivity and chemoselectivity has become a much challenging and demanding area for synthetic chemists. In this respect, we have developed **diastereoselective synthetic protocols** for 1,3-thiazines **5** (Ref. 5) and azolopyrimidines **6** (*Tetrahedron* **2006**, 62, 5464) using 2-methyl-2-phenyl-1,3-oxathiolan-5-one, a novel mercaptoacetyl transfer agent **7** reported from our laboratory (*Tetrahedron* **2005**, 61, 10013). In addition, we have also developed a **diastereoselective green protocol** for annulation of *s*-triazine ring on thiazoles **8** (*Green Chem.* **2006**, 8, 455) and have devised a **chemoselective annulation** of 1,3-dithiin, -thiazine and -oxathiin rings on thiazoles **9**, **10**, and **11** involving tandem Knoevenagel, Michael and ring transformation reactions employing solvent-free MW irradiation conditions in a one-pot procedure (*Tetrahedron* **2006**, 62, 8029). Very recently, **regio/- and stereoselective synthetic routes** for amino/ mercaptodithiazolopyrimidines **12** from thiourea have been developed by us (*Tetrahedron* **2007**, 63, 6924, *Synthesis* **2007**, 3831).



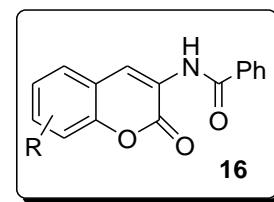
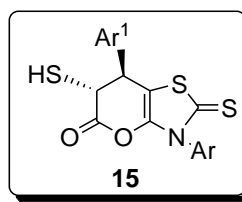
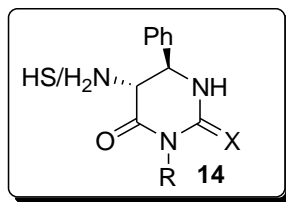
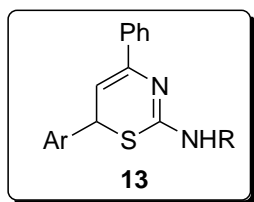
### [D]. Active-Copper Promoted *N*-Arylations in Aqueous Media:

We have reported active-copper-promoted mild and expeditious *N*-arylations of amines, amides, imides, and  $\beta$ -lactams with aryl halides under MW irradiation conditions. These reactions can be performed in aqueous media as well as under solvent-free conditions to give good yields (*Synthesis* **2006**, 1868). Interestingly, no base is used in these reactions and the active copper itself acts as the halogen acceptor. Very recently, we have reported the catalyst-free synthesis of pharmaceutically and chemically important 3-mercaptocoumarins in water (*Green Chem.* **2009**, 11, 878).

### [E]. Ionic Liquid Mediated Organic Synthesis:

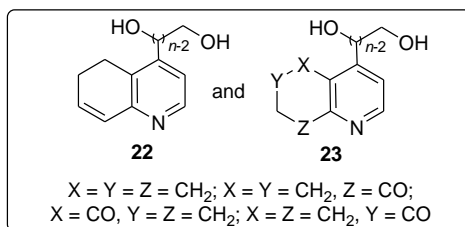
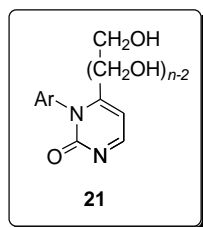
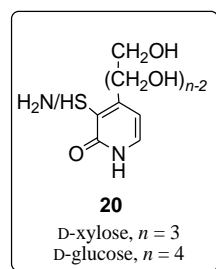
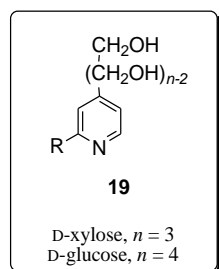
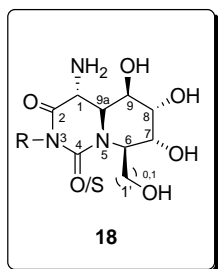
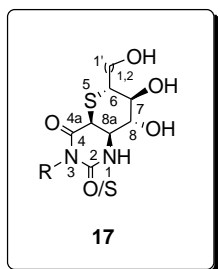
We have disclosed an efficient conjugate hydrothiocyanation of chalcones with a *task-specific ionic liquid* ([Bmim]SCN). The application of this protocol in heterocyclic chemistry is demonstrated by a one-pot synthesis of 1,3-thiazines **13** (*Tetrahedron Lett.* **2007**, 48, 7793). A chiral ionic liquid-catalyzed, efficient and unprecedented version of the Biginelli reaction using novel variants of its active methylene component, viz. 2-phenyl-1,3-oxazol-5-one/2-methyl-2-phenyl-1,3-oxathiolan-5-one, with benzaldehyde and urea/thiourea enantio- and diastereoselectively, yields 5-amino-/mercaptoperhydropyrimidines **14** (*Tetrahedron* **2008**, 64, 1420). Recently, a stereocontrolled route to mercaptopyranothiazole **15** has been developed using one-pot multi-component protocol in chiral ionic liquid (*J. Heterocyclic Chem.* **2008**, 45, 1315). Furthermore, we have also developed a one-pot protocol for 2-amino-5-mercapto or 2,5-diamino-1,3-thiazines using ionic-liquid ([Bmim]Br) (*Tetrahedron* **2009**, 65, 1306). Very recently, we have reported the one-pot [Bmim]OH-mediated synthesis of 3-benzamidocoumarins **16** (*Tetrahedron Lett.* **2009**, 50, 2208).





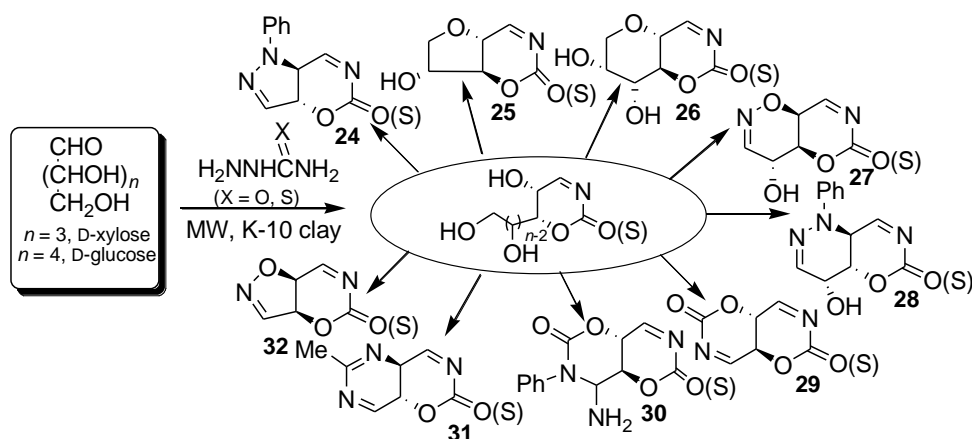
### [E]. Biorenewable Resources in Organic Synthesis:

“A raw material as feedstock should be renewable rather than depleting wherever technically and economically practicable.” This quotation is one of the 12 principles of green chemistry and thus, “renewable resources” is a new and rapidly developing concept in the environmental and chemical sciences that concerns the wide use of biorenewable materials for industry. In this context, we have utilized carbohydrate feedstocks as raw material in “**Biginelli Reaction**” and we have developed an efficient Ce(III)-catalyzed diastereoselective synthesis of iminosugar annulated-(**17**, *Synlett* **2007**, 1905) and montmorillonite K-10 clay-catalyzed annulated (**18**, *Tetrahedron Lett.* **2007**, 48, 4899) polyfunctionalized pyrimidine scaffolds of pharmacological potential under solvent-free MW irradiation conditions in a one-pot procedure. Recently, we have developed an original method for polyhydroxyalkylpyridines (**19**) and their 3-amino/mercapto-2-pyridinone analogues (**20**) using unprotected aldoses as biorenewable resources via enol-driven Michael-type addition of lactones/ketones to aldose-derived 1,3-oxazin-2-ones followed by decarboxylative ring transformation (*Synlett* **2008**, 529). Very recently, we envisaged a K-10 clay-catalyzed amine-driven dehydrative ring transformation approach to pyrimidines (**21**, *Tetrahedron Lett.* **2008**, 49, 2377) and enol-driven ring transformation approach to dihydro- (**22**, *Synlett* **2008**, 2257) and tetrahydroquinolines (**23**, *Synlett* **2008**, 2257) from carbohydrates as biorenewable resources.



### [F]. Diversity-Oriented Synthesis:

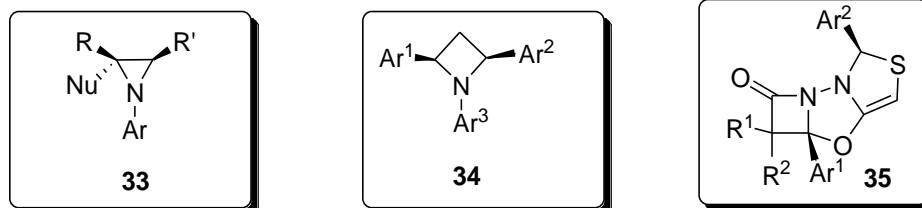
We have developed a general, straightforward diversity-oriented synthetic approach for the synthesis of various 1,3-oxazin-2-one(thione)-fused *N*- and *O*-heterocyclic systems (**24-32**) using D-glucose and D-xylose as biorenewable resources under solvent-free microwave irradiation conditions (*Tetrahedron* **2008**, 64, 4246) and also by employing  $\text{Cu}(\text{OTf})_2$  as catalyst (*J. Chem. Res.* **2009**, 522) where,  $\text{Cu}(\text{OTf})_2$  acts as dehydrazinating and *N*-aryllating reagent.



### [G]. Synthesis of Small Ring N-Heterocycles:

Owing to the inherent strain in small ring heterocycles, they are useful as feedstocks in organic synthesis to provide functionalized carbon chain. A general method for a convenient synthesis of 1,2,4-trisubstituted azetidines **34** by reductive cyclization of readily available aza-Michael adducts of chalcones and diethyl *N*-arylphosphoramidates in a one-pot procedure is reported (*Tetrahedron Lett.* **2007**, 48, 8037), which may find application in organic synthesis. Furthermore, we have developed a novel and efficient aziridination of  $\alpha$ -halo ketones (*Tetrahedron Lett.* **2008**, 49, 687). The reaction of  $\alpha$ -halo ketones with diethyl *N*-arylphosphoramidates affords diethyl *N*-aryl-*N*-(2-oxoalkyl)phosphoramidates which undergo reductive ( $H^-$ -induced) cyclization with sodium borohydride followed by sodium hydride to give 1,2-disubstituted and 1,2,3-trisubstituted aziridines. The cyclization induced by  $NCS^-$  or  $PhS^-$  affords substituted aziridines functionalized at position 2. The reactions give excellent yields and are highly diastereoselective in favour of *cis* aziridines **33**.

Recently, tricyclic  $\beta$ -lactam antibiotics, generally referred to as “*trinems*”, have been the subject of considerable study owing to their broad spectrum of antibacterial activity. Besides, the ever-growing new applications of 2-azetidine in fields ranging from enzyme inhibition to the use of these products as starting materials to develop new synthetic methodologies have triggered a renewed interest in the building of new polycyclic  $\beta$ -lactam systems in an attempt to move away from the classical  $\beta$ -lactam antibiotic structures. In this regard, we envisaged the straightforward synthesis of trinem class of antibiotics, viz highly derivatized azetidino[2,1-*b*]-thiazolo[3,4-*d*]-3*H*-1,3,4-oxadiazol-6-ones **35** (*Tetrahedron Lett.* **2008**, 49, 5553).



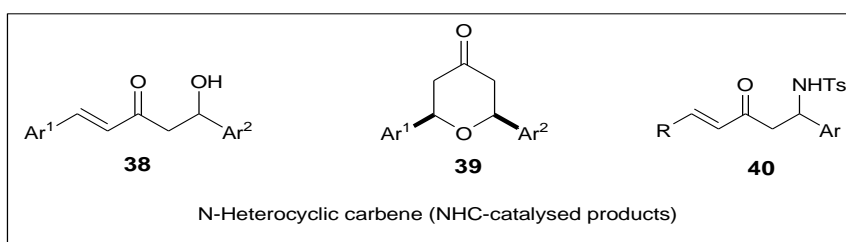
### [I]. Morita-Baylis-Hillman (MBH) Chemistry:

Morita-Baylis-Hillman (MBH) adducts bearing allylic hydroxyl and Michael acceptor units and their derivatives have been illustrated as valuable synthons and starting materials for the generation of diverse molecular skeletons employing simple alternatives.<sup>1</sup> Especially, the regioselective introduction of nucleophiles at either  $\alpha$ - or  $\gamma$ - position of the MBH acetates enables the construction of a variety of bioactive molecules and has become a powerful tool in synthetic organic chemistry. Thus, we have performed the first regio- and diastereoselective direct introduction of  $\alpha$ -mercaptoacetic acid/amide units into MBH acetates **36** (*Tetrahedron Lett.* **2009**, 50, 2414). Furthermore, we have also reported a novel one-pot stereoselective synthesis of *N*-protected  $\alpha$ -Amino acids from MBH Acetates **37** (*Synlett* **2009**, 1423).



### [J]. N-Heterocyclic Carbenes (NHCs)-catalyzed organic synthesis:

Over the last decade, there have been a continuously growing number of successful and novel applications of *N*-heterocyclic carbenes (NHCs) as organocatalysts and reagents for an expanding set of reactions. This is not only because of the great versatility of these organocatalytic transformations but also due to the possibilities that arise from the NHC's characteristic causing an inversion of the classical reactivity, that is, umpolung. We have disclosed an NHC-catalyzed efficient synthesis of aldol products  $\beta$ -hydroxy- $\alpha,\beta$ -unsaturated ketones **38** via carbonyl umpolung reaction of enals with terminal epoxides. Furthermore, we also demonstrated its synthetic application and developed a straightforward, convenient, and one-pot process for the synthesis of tetrahydropyran-4-ones **39** in excellent yields (87-90%) via oxy-Michael intramolecular reaction of **38** (Synlett 2010, 240-246). Recently, we have reported an unprecedented synthesis of  $\beta'$ -amino- $\alpha,\beta$ -unsaturated ketones



**40** via regioselective aziridine ring-opening. The protocol involves carbonyl umpolung reactivity of enals in which the carbonyl carbon attacks nucleophilically on electrophilic terminal aziridines regioselectively (Tetrahedron Lett., 2010, 51, 1657).

### [H]. Synthesis of Graphene-based materials:

We have prepared Fe<sub>3</sub>O<sub>4</sub> decorated Naf/DMAP linked graphene oxide (Fe<sub>3</sub>O<sub>4</sub>-Naf/DMAP-GO) (Fig. 41) nanohybrid film which worked well as a highly selective non-enzymatic electrochemical sensor. The linking of 3,7-bis (Dimethylamino)-phenothiazine-5-ium chloride (DMAP) occurs via electrostatic interaction of the cationic organic compound with negatively charged oxygen-containing groups (-COO<sup>-</sup> and -O<sup>-</sup>) available on the edge of graphene oxide (RSC Advances, 2016, 6, 104868). Recently we have reported facile construction of AuNPs modulated SDS wrapped G-TC tailored electrode (Fig. 42) for sensitive detection of ascorbic acid (New J. Chem. 2017, 41, 6489).

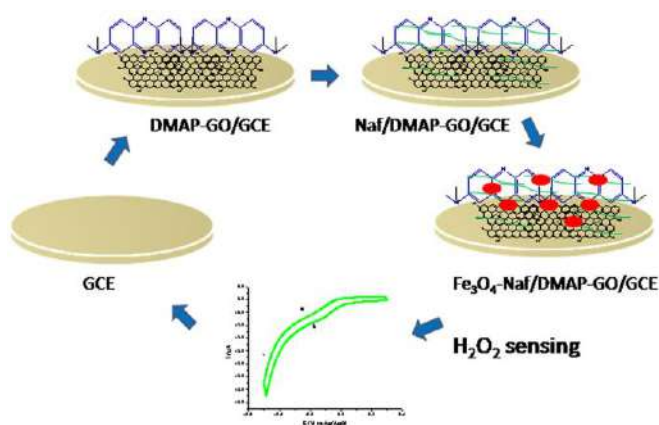


Fig. 41: Preparation of Fe<sub>3</sub>O<sub>4</sub> decorated GO.

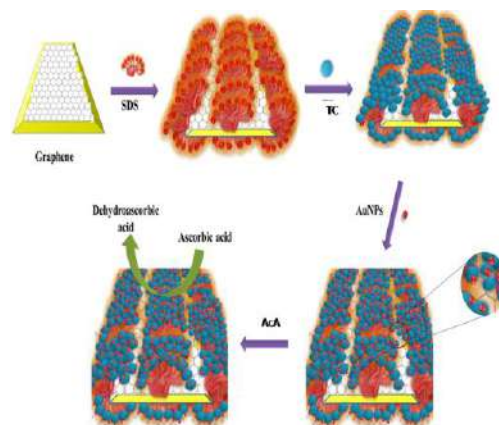
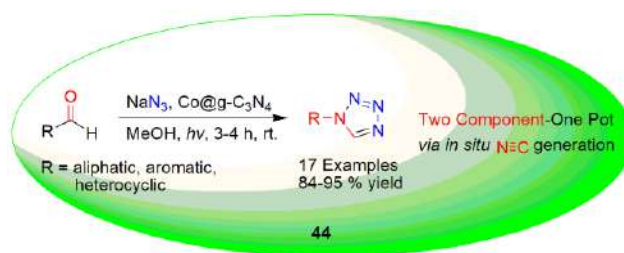
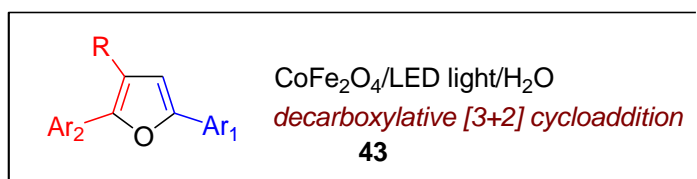


Fig. 42: Preparation of Au-decorated GO.

## [I]. Synthesis of Nano-materials and Photo-catalysis:

We are involved in the synthesis of nanomaterials and their application in photo-catalysis. We have reported the synthesis of  $\text{CoFe}_2\text{O}_4$  nanoparticles and their catalytic behaviour as its co-operative effect along with visible light photo-catalysis for green synthesis of furans in water (Fig. 43). Though, the reported method is efficient without catalyst in presence of visible light (70% yield in 4h at rt), the use of catalyst not only increases the yield (91%) but also accelerates the conversion rate (2h, rt) (*New J. Chem.* **2017**, *41*, 4937). Recently, a novel and green  $\text{Co@g-C}_3\text{N}_4$  catalyzed visible-light-driven direct regioselective synthesis of 1*H*-tetrazoles (**44**) directly from various aldehydes and sodium azide is reported (*Green Chem.* **2018**, DOI: 10.1039/C8GC01321G). Herein,  $\text{NaN}_3$  not only behaves as a three-nitrogen donor of the tetrazole ring but also converts aldehyde into isocyanide as the one-nitrogen source.



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