Review Paper Aedes Mosquito Dynamics: Unravelling Behavior, Genetics, and Arbovirus Risks in India



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ABSTRACT

Background: Aedes aegypti and Aedes albopictus responsible for transmitting severe viruses, such as dengue, chikungunya, yellow fever, and Zika, pose a significant health threat in India. Challenges in vector control, including insecticide resistance and limited progress in vaccine development, coupled with factors like climate change and urbanization, contribute to an extensive expansion of mosquito habitats and disease transmission. However, the existing literature on how these factors influence mosquito behavior, genetic architecture, and virus transmission patterns in different regions of India is insufficient.

Methods: This scoping review conducted a thorough search across online databases, such as Google Scholar, Web of Science, Scopus, and PubMed. The keywords that were used were as follows: "*Ae. Aegypti*," "*Ae. albopictus*," "breeding habitat," "feeding preference," "genetic diversity," "virus transmission," "dengue," "chikungunya," "Zika," and "India." The online search, conducted from October to November 2023, focused on English-language articles from 2013 to November 2023. The inclusion criteria encompassed articles addressing *Aedes* behavior, genetic diversity, and disease transmission, while the exclusion criteria removed non-*Aedes* mosquito studies and duplicates. Article quality was manually checked and assessed.

Results: The literature survey spanning the past ten years categorized the data into the following four subgroups: "Arboviral disease," "arbovirus transmission," "*Aedes* mosquito behavior," and "genetic diversity." The data synthesis emphasized the need to focus on *Aedes* mosquito breeding habitats and virus transmission patterns in common dengue endemic regions, like Delhi, Tamil Nadu, Kerala, Rajasthan, and Uttar Pradesh in India. However, research on feeding behavior and genetic diversity is scarce. Hence, more research should be conducted in these areas.

Conclusion: This analysis highlights critical gaps in our understanding, emphasizing the necessity for detailed mapping, studying virus evolution, socio-epidemiological research, and using advanced technologies for real-time monitoring. Understanding the complex relationships between blood-feeding patterns, host preferences, and diverse breeding habitats is crucial for effective disease control.

Keywords: Dengue, Aedes, Behavior, Genetic diversity, Virus transmission

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Introduction

engue is a predominant arboviral disease transmitted by *Aedes* (Stegomyia) *aegypti* and *Aedes* (Stegomyia) *albopictus* mosquitoes, pose a significant global threat, contributing to an annual burden of 100–400 million new infections. Approximately 3.83 billion people are at risk worldwide [1, 2].

These vectors also transmit other impactful diseases, such as chikungunya, yellow fever, and Zika infections [3]. The spectrum of dengue infections encompasses mild fever to severe forms, like dengue hemorrhagic fever and dengue shock syndrome, driven by four distinct serotypes (DENV– 1, DENV–2, DENV–3, and DENV–4) capable of co-infecting humans [3, 4]. Dengue clinical manifestations include fever, rash, eye pain, arthralgias, myalgias, and hemorrhage [3].

The quantification of the global burden of dengue is challenging due to widespread under-reporting, especially in areas with weaker healthcare systems [5, 6]. Dengue is widespread, affecting over 120 countries, especially in the Americas, Southeast Asia, and the Western Pacific [5, 6]. The global spread of dengue began in the 1980s, reaching various regions, such as the Mediterranean basin, the Americas, and Africa [7]. Brazil has the highest number of reported cases in the Americas, with 4.1 million, while Africa, particularly Burkina Faso, is experiencing a significant increase in the number of cases [7, 8]. Dengue is also reported in Eastern Mediterranean countries, with the emergence of local cases and the expansion of Ae. albopictus in Europe [9]. In the Western Pacific Region, which includes Australia, China, and the Philippines, there are over 500000 reported cases [7]. About 70% of the disease burden is concentrated in Asian countries [2, 5, 6]. South-East Asian nations, like India, Indonesia, Myanmar, Sri Lanka, and Thailand rank among the top 30 countries globally with the highest dengue prevalence, as per the Centers for Disease Control and Prevention 2024 [4, 10].

India's contribution to global dengue infections is estimated at 34 out of 96 million, significantly contrasting with the reported 12 484 cases to the World Health Organization (WHO), revealing a substantial underreporting gap [2]. However, in India, there is a lack of clear understanding of the distribution of dengue vectors, although their presence has been confirmed in specific regions through local surveillance efforts [11-13]. Major regions in India, are affected by one of the four arboviral infections [14]. Furthermore, a study on the impact of climate change on mosquito abundance in India suggests an increased risk in parts of the Thar Desert and the Upper Himalayas [15]. Despite lower reported dengue cases, there is a high prevalence observed in northeast India and the western coastline, emphasizing the need for heightened monitoring [11]. *Ae. aegypti* dominates in the Deccan plateau and arid regions of Gujarat and Rajasthan, while *Ae. albopictus* prevails in the eastern coastline, aligning with reported dengue incidence in Gujarat, Maharashtra, Punjab, and Karnataka [11, 15].

Ae. aegypti and Ae. albopictus showcase adaptability shaped by their evolutionary history and genetic diversity [9, 16]. Ae. aegypti has two subspecies, namely Ae. aegypti aegypti and the ancestral Ae. aegypti formosus, originating from Africa, with the former adapting to urban settings and becoming a primary vector [16]. Ae. albopictus, from Southeast Asia, displays versatility in human-altered environments, spreading worldwide [8]. Behaviorally, Ae. aegypti prefers indoor resting and daytime human feeding, mainly in urban areas and artificial water containers [3, 4]. Conversely, Ae. albopictus, an opportunistic feeder with outdoor resting tendencies, also exhibits anthropophagic behavior [1, 4]. Factors influencing their distribution include urbanization, climate elements (temperature, rainfall, and humidity), and human mobility [1, 4, 6, 11, 17]. Human movement contributes to lower genetic differentiation [15], while changing precipitation patterns impact larval habitats, affecting vector population dynamics [16]. The intricate interplay of these factors contributes to the rapid expansion of vectors and their prevalence worldwide [1, 4, 6, 8, 9, 11, 17].

Despite the increasing number of dengue cases, there is no specific treatment or vaccine available [2, 7, 10, 14]. In India, dengue control relies on different practices, such as indoor spraying, fogging, and personal protection promotion, yet strong evidence of the effectiveness of these interventions is lacking [18, 19]. This underscores the urgent need for tailored control measures. Conventional surveillance often overlooks factors, like seasonal larval abundance fluctuations and non-residential areas within cities, where Aedes mosquitoes thrive [20]. The impact of urbanization, social behavior, and climate change on Aedes mosquitoes in India are not well understood. Recent studies highlight challenges in controlling Ae. aegypti and Ae. albopictus in Western Africa and Thailand due to insufficient research on their behavior [1, 4]. Recognizing the global impact of Aedes-borne diseases, our study addresses knowledge gaps in Aedes mosquito behavior, genetic diversity, and arbovirus transmission in India [1, 4]. To enhance surveillance and guide control efforts, this study pinpoints research gaps in Aedes mosquito behavior in different regions of residential areas and non-residential areas [11, 17, 20].

Methods

This study employed a scoping review approach by following the methodology from [21]. The article selection process involved five steps as follows: Defining the research question, conducting a thorough search for relevant studies, filtering and selecting pertinent articles, synthesizing and interpreting collected data, finally establishing the analytical framework, and reporting the results. Searches were conducted in Google Scholar, Web of Science, Scopus, and PubMed from October to November 2023, focusing on articles published between 2013 and November 2023. We have subdivided the second step of our search strategy into three substep processes: 1) An initial search of published studies in Google Scholar, Scopus, PubMed, and Web of Science databases; 2) Utilization of keywords and terms to search additional databases, including Science Direct, relevant government ministry portals (e.g. National Center for Vector Borne Diseases Control (NCVBDC)) and organizations (e.g. WHO and centers for disease control and prevention) by using Google search engine and manual search; 3) Exploration of the reference lists of identified studies for supplementary sources. Appropriate keywords were used to search literature in English language only and articles in other languages were removed. Whenever feasible, we used the preferred reporting items for systematic reviews and meta-analysis criteria [22] for performing and writing the results (Figure 1). The keywords used to search literature are as follows:

• "Ae. aegypti" AND "dengue" AND "India". "Aedes albopictus" AND "dengue" AND "India"

• "Ae. aegypti" AND "chikungunya" AND "India". "Ae. albopictus" AND "chikungunya" AND "India"

• "Ae. aegypti" AND "Zika" AND "India". "Ae. albopictus" AND "Zika" AND "India"

• "Ae. aegypti" AND "breeding" or "breeding habitat" AND "India". "Ae. albopictus" AND "breeding" AND "India"

• "Ae. aegypti" AND "feeding preference" or "feeding behavior" or "blood-feeding" AND "India". "Ae. albopictus" AND "feeding preference" or "feeding behavior" or "blood-feeding" AND "India"

• "Ae. aegypti" AND "genetic diversity" or "genetic structure" or "genetic variability" or AND "India". "Ae. albopictus" AND "genetic diversity" or "genetic structure" or "genetic variability" or AND "India"

• "Aedes mosquitoes" AND "virus transmission" or "vertical transmission" or "transovarial transmission" AND "India"

The inclusion criteria for this review article were limited to full articles in the English language, concentrating on the status of Aedes borne diseases in India, Aedes mosquito behavior, including feeding behavior, breeding habitats, genetic diversity, arbovirus transmission dynamics in mosquitoes, and patterns of transmission in hosts. The review considered the influence of external factors, like ecological elements (including rainfall, temperature, and humidity) and social factors of urbanization, on mosquito adaptation to changing environments and their increased expansion. Data collection involved research on Ae. aegypti and Ae. albopictus across diverse states in India. The quality of the articles was checked and stored manually. Meanwhile, the exclusion criteria targeted articles that did not focus on the behavior and genetic diversity of Aedes mosquitoes in the Indian context, articles related to mosquitoes other than Aedes and the removal of duplicated or commonly addressed articles. The screening criteria for selecting the objectiverelated articles included having relevance to the study subject.

Results

Following an initial search and review of titles and abstracts, we identified a total of 102 articles from the online search and an additional 4 articles through manual searching. The details of these articles are outlined in Table 1. To align with the study objective, the extracted data from these articles were categorized into specific themes, including "*Aedes* mosquito biology and arboviral disease in India," with sub-categories, such as "arboviral disease in India," "virus transmission dynamics," "feeding behavior, host preference, and biting pattern," "breeding habitat," and "genetic diversity."

Arboviral disease in India

In the past two decades, India has witnessed a substantial geographic spread of dengue, experiencing repeated outbreaks and a staggering 11-fold surge in cases [23]. The global dengue outbreak in 2016, coupled with Delhi's severe outbreak in 2015, underscored the widespread impact of the virus [23]. Dengue's presence in India dates back to 1950, with the isolation of the first Dengue serotype, DENV-1, in Vellore, Tamilnadu, in 1956, followed by a major outbreak in Calcutta in 1963 [23]. Since the 1960s, outbreaks have been reported across all states and union territories, except Ladakh and Lakshad-



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Figure 1. Flowchart of article selection process adopted from preferred reporting items for systematic reviews and meta-analyses [22]

weep [23]. The emergence of dengue hemorrhagic fever intensified outbreaks, particularly in Delhi in 1996 and North India in 2003, highlighting the virus's variability [24, 25]. According to the NCVBDC, data as of January 2023 (Figures 2A, 2B, and 2C) depict the distribution of reported dengue cases and deaths in India. Subsequent reports in September 2023 indicate a sustained impact of dengue fever, with persistently high cases in states like Kerala, Karnataka, Maharashtra, Odisha, Uttar Pradesh, Assam, Delhi, Telangana, Rajasthan, Tamil Nadu, West Bengal, and Punjab [14, 26].

A nationally representative survey in India unveiled age-specific dengue seroprevalence, estimating an overall prevalence of 48.7%. Variations were observed, ranging from 28.3% in children aged 5–8 years to 56.2% in individuals aged 18–45 years [27]. Geographical differences were evident, with the highest seroprevalence in southern regions (76.9%) and urban areas (70.9%), underscoring the impact of urbanization on dengue incidence [27]. Furthermore, genomic analyses reveal the co-circulation of all four dengue serotypes in different regions of India, with DENV-4 emerging as a dominant serotype in South India [28]. Diagnosis involves detecting NS1 antigen and IgM dengue antibody. Vaccines like TAK-003 are undergoing trials, facing challenges due to diverse antigens and genotype variations [29].

Chikungunya fever in India, caused by Chikungunya virus (CHIKV), is endemic in 24 states and 6 union territories [30]. The virus reemerged in 2005, causing a significant outbreak across states, with CHIKV's adaptability to Ae. albopictus mosquitoes contributing to its rapid spread [31]. Molecular diagnostics, particularly reverse transcription polymerase chain reaction, play a crucial role in diagnosing acute-phase CHIKV infections [30]. Additionally, the Zika virus (ZIKV), with a complex evolutionary history, surfaced in Gujarat and Maharashtra in 1954, reappearing notably in Ahmedabad in 2016, Jaipur in 2018, and Madhya Pradesh in 2021 [32-34]. The continuous emergence of these viruses underscores the importance of robust surveillance, prompt diagnosis, and effective prevention strategies tailored to regionspecific strains [33, 34].



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Figure 2. Dengue cases and death reported in India (2019-2023 continuation as per NCVBDC in India) [14]

A) Various states and union territories in India impacted by dengue fever, B) Total number of reported dengue cases and associated deaths from 2019 to January 2023, C) Distribution of dengue cases and death cases across different states and union territories in India.



Figure 3. Types of virus transmission observed in *Aedes* mosquitoes [3] **WRP** Notes: Green represents horizontal transmission, pink represents vertical transmission, and blue represents venereal transmission.

Virus transmission dynamics

Arboviruses, transmitted primarily by mosquitoes, utilize horizontal and vertical transmission mechanisms. Horizontal transmission occurs during a blood meal, while vertical transmission involves passing the virus from an infected female mosquito to her offspring through eggs (Figure 3) [35, 36]. Understanding these mechanisms is crucial for studying dengue fever epidemiology. Global studies, including in India, confirm natural vertical transmission of dengue virus in *Aedes* mosquitoes [37]. Indian studies since 2007 show natural vertical transmission of dengue serotypes, varying globally, with South America and Asia contributing significantly to cases [35]. Africa lacks sufficient research on natural vertical transmission [38].

In India, studies in various states have shed light on the intricate dynamics of arbovirus transmission, particularly in Chennai, emphasizing the role of vertical transmission in maintaining dengue viruses during interepidemic periods [39]. Kerala reports natural vertical transmission in *Ae. albopictus* [40]. Rajasthan investigates dengue-3 virus persistence in successive *Ae. aegypti* generations, emphasizing re-emergence [41]. Uttar Pradesh reveals natural vertical transmission of dengue virus seen in male *Aedes* mosquitoes [42]. Delhi's 2013 outbreak shows *Ae. aegypti* infection rates contribute to transmission in low and medium-income areas [43]. In Jodhpur, Bhopal, *Aedes* mosquitoes tested positive for dengue antigen via vertical transmission [44, 45]. Odisha faced rising dengue cases, prompting studies on DENV transmission dynamics [46].

Furthermore, vector surveillance is crucial, as seen in central India's year-round xenomonitoring [47]. Nationwide surveillance initiated by the Indian Council of Medical Research from 2016 to 2019 across 14 states contributed to our understanding of ZIKV and DENV prevalence in *Aedes* mosquitoes [48]. Thiruvananthapuram's intensive vector surveillance during a ZIKV outbreak detected high *Aedes* mosquito infestation [33].

Feeding behavior, host preference, and biting pattern

Hematophagy, essential for arthropod vector life cycles, underscores the frequent blood feeding of disease vectors like *Ae. aegypti* and *Ae. albopictus*, occurring every 2 to 4 days, is crucial for pathogen acquisition and transmission [49]. These mosquitoes exhibit a strong preference for human blood, influenced by environmental factors and host availability [50, 51]. In Asia, where *Aedes* mosquitoes are prevalent, they serve as competent vectors for various infectious diseases [2]. The body size of these mosquitoes, influenced by temperature and nu-

Behavior	A. aegypti			A. albopictus		
	Major Findings	States	Gap in Knowl- edge	Major Findings	States	Gap in Knowl- edge
Host seeking behavior	Displays a diurnal biting pattern, aligning with its role as a primary vector of dengue and Chikungunya in India [51, 52, 54].			Biting rhythms of <i>Aedes</i> mos- quitoes reveal a diurnal pattern [51, 52]		
Breeding habits	Favor indoor breeding within households; key containers, like plastic contain- ers, overhead tanks, in <i>Aedes</i> breeding dynamics [59], discarded auto- mobile tires, and water-holding plastic contain- ers, for breeding [60, 67]	Delhi [67, 94, 72, 98], Kerala [52-54, 71, 99], South Anda- man [73, 100], Banguluru [17], Odisha [59, 60], Tamilnadu [61, 62], Assam [101], Udaipur, Rajast- han [102], Thiru- vanantapuram [71], Bihar [67], Dehradun [76]	Limited informa- tion on the adaptability of <i>Aedes</i> mosqui- toes to diverse breeding habi- tats in different regions of India [51, 64]	Shows a prefer- ence for outdoor areas, such as car tires and garbage dumps, discarded auto- mobile tires and water-holding plastic contain- ers [59], rubber collection bowls and various artificial contain- ers [60]	Tripura [69], Kerala [64, 65], Thrissur, kanyakumari [70, 68], Thiru- vanantapuram [71], Banguluru [20], Assam [101], Tamilnadu [61, 62], Odisha [59, 60] and Bihar [67], Dehradun [76], West Ben- gal [72, 91]	Limited informa- tion on the spe- cific ecological and environ- mental factors influencing the adaptability of <i>Ae. albopictus</i> to diverse breeding preferences [77, 61]
Genetic diversity	High level of genetic diversity among <i>Ae. ae- gypti</i> popula- tions, indicating human-medi- ated dispersal and connectivity within localized regions [95]	South India [78], Delhi 86], Northeast region [87], Rajasthan [89]	Limited genetic studies com- pared to global research efforts [95]; further re- search is needed to explore the mechanisms and frequency of mosquito move- ment through transportation networks and migration routes in India [95]	Low genetic diversity globally due to factors like insecticide use, ecological adaptability, and human-mediat- ed transporta- tion [103]; rapid global dispersion driven by inter- national trade and environ- mental changes [62]	Odisha [92], Assam [88], Tamilnadu [85], Thiruvanantha- puram [79]	Limited informa- tion on the spe- cific ecological and environ- mental factors influencing the adaptability of <i>Ae. albopictus</i> to diverse breeding preferences [95]
Virus transmis- sion	Horizontal and vertical ma- jorly transovar- ial transmission mechanisms arbovirus transmission observed [35]; natural vertical transmission of dengue virus in <i>Ae. aegypti</i> in India [36]	Chennai [39], Kerala [40], Rajasthan [41], Uttar Pradesh [42], Delhi [43], Jodhpur [44], Bhopal [45], Odisha [46]	Limited un- derstanding of the molecular mechanisms of vertical and transovarial transmission. [40]; need for deeper research considering the evolutionary as- pects of the <i>Ae-</i> <i>des</i> mosquito's multiple feeding behaviors	Horizontal and vertical transmission mechanisms play crucial roles in arbovirus transmission in <i>Ae. albopictus</i>	Kerala [40], Odisha [46]	Limited un- derstanding of the molecular mechanisms of vertical and transovarial transmission [36]

Table 1. Summary of the major findings on Aedes mosquito behavior, genetic diversity, and virus transmission dynamics in India



Figure 4. A conceptual framework representing the factors contributing to the escalation of *Aedes* mosquito populations and proposing strategies to address the prevalence of dengue [1, 4, 6]

trition, impacts their ecology and multiple blood-feeding activities [52]. A study in the South Andaman district found that both *Ae. aegypti* and *Ae. albopictus* predominantly feeds on humans, with *Ae. albopictus* exhibiting a broader host range [53]. In Dehradun City, variations in mosquito abundance based on different habitats and seasons were observed, with *Aedes* mosquitoes primarily feed on humans, followed by bovines and pigs [54]. Understanding biting rhythms is also crucial, as evidenced by a study in Sivaganga district identifying three main rhythmic patterns among various mosquito species [55]. Additionally, the innovative approach of using attractive toxic sugar baits in India has shown efficacy in reducing mosquito populations by exploiting the sugar-feeding behavior of both male and female mosquitoes [56, 57].

Breeding habitat

India's diverse landscapes and climates create a complex scenario for *Aedes* mosquito breeding, key vectors for diseases like dengue [58]. *Aedes* mosquitoes are adaptable to various habitats, from clean water receptacles to polluted collections like stagnant drain water, tanks, coolers, and containers [59]. *Ae. aegypti* favors indoor breeding, while *Ae. albopictus* prefers outdoor areas, like car tires and garbage dumps [59]. Visual and olfactory cues guide their oviposition site selection, considering both chemical and physical factors in the water [58]. Extensive studies across India emphasize *Aedes* mosquitoes' adaptability to diverse breeding habitats, with larval indices, like house index, container index, and breteau index, serving as crucial tools for assessing dengue epidemiological risk [60]. In Delhi and Tamil Nadu, significant *Aedes* breeding dynamics occur in key containers like overhead tanks, enhancing cost-effective and sustainable control efforts [61, 62].

Kerala, susceptible to arboviral diseases, reveals unconventional breeding habitats in crops and brackish water areas [63, 64]. Northern Kerala studies link physicochemical parameters with adult emergence, providing insights into the relationship between water quality and mosquito presence [65]. The adaptability of Ae. albopictus to diverse environments, including discarded automobile tires and water-holding plastic containers have been observed in the Andaman Islands [66]. Various regions, such as Kanyakumari, Tripura, Thrissur, Bengaluru, Odisha, Bihar, Dehradun, and West Bengal identify specific containers, like plastic containers, tires, and water coolers as high-risk areas for Aedes breeding [20, 60, 67-77]. Entomological surveillance international airports underscore varying degrees of receptivity to Aedes vector breeding, emphasizing the importance of consistent monitoring and timely source reduction efforts [76].

Genetic diversity

In India, rising dengue cases necessitate a multidisciplinary approach, integrating morphological, molecular, and distributional data [78]. Our data analysis focuses on two key aspects as follows: First, understanding the origin and evolution of dengue serotypes, and exploring how they adapt to changing environments is one aspect. Secondly, we delve into the genetic variation within Indian Aedes mosquito populations compared to those in other countries. The origin and evolution of the dengue virus is complex, and not as clear as of some other viruses. Cases resembling dengue were recorded in the early 1600s in the Caribbean and Panama, with unambiguous descriptions emerging in Africa, Asia, and the Americas in 1779–1788 [79]. DENV was present in Asia in the 1700s before the arrival of Ae. aegypti, suggesting transmission by another mosquito, likely Ae. albopictus [79]. The prevailing belief is that the current four serotypes of the dengue virus originated in Asia. Recent evidence suggests that over 300 years ago, ancestral dengue virus likely existed in Africa [79]. Furthermore, phylogenetic relationships with other flaviviruses, indicate the dengue virus's affiliation with African lineages, emphasizing a historical transmission by African mosquitoes to primates [79].

Given the significant public health impact of dengue infections in India, understanding genetic diversity, spatial distribution, vaccine effectiveness, and potential new variants is crucial [80]. All four dengue serotypes (DENV-1-4) have been isolated from various parts of India, with DENV-2 and DENV-3 recognized as the most dominant genotypes in terms of disease spread, while DENV-1 has been predominantly observed in recent decades [80]. The genetic diversity and antigenicity of the dengue virus primarily depend on mutations in its envelope protein [80]. Phylogenetic analysis of the envelope protein further subdivides these serotypes into distinct genotypes [81]. A study involving 50 Indian strains, including 19 recombinant strains, indicated that DENV-3 exhibits less sequence diversity compared to other serotypes [81]. Positive selection on several codons correlated with genetic diversity between serotypes, highlighting that amino acid diversity and inter-genotypic recombination are major contributors to antigenicity variation and dengue virus evolution within India [81].

Global studies on *Ae. aegypti* reveals two primary genetic units: *Ae. aegypti formosus* in Africa and *Ae. aegypti aegypti* outside Africa [82]. In contrast, *Ae. albopictus*, due to insecticide use, adaptability, and human transport, shows limited genetic diversity, spreading rapidly worldwide [83]. Molecular markers like isozymes, restriction fragment length polymorphism, relative afferent pupillary defect, mitochondrial DNA, ribosomal DNA, and microsatellites [83, 84] help to understand genetic diversity in disease-carrying mosquitoes [85]. South India identifies Ae. aegypti haplotypes, revealing global genetic differences [78]. In Delhi, relative afferent pupillary defect analysis shows significant genetic variation linked to urbanization and climate conditions [86]. North-east regions explore genetic diversity and unique Wolbachia infection patterns in Ae. albopictus [87]. Sonitpur district in Assam uses DNA barcoding, unveiling notable Ae. albopictus nucleotide variations [88]. Maharashtra examines COI markers, revealing low genetic variability in Ae. aegypti populations [84]. Rajasthan, using ITS-2 and mitochondrial COI markers, shows low genetic variability in Ae. aegypti's COI gene [89]. Tamil Nadu's COX1-based DNA barcoding effectively identifies Ae. albopictus, emphasizing the importance of genetic polymorphism awareness [79]. Comprehensive genetic structure analysis in 22 Indian populations indicates high genetic diversity among Ae. aegypti populations, influenced by human-mediated dispersal and connectivity [17]. Genetic diversity and evolutionary adaptation in mosquitoes are crucial for predicting their future survival, development, and migration patterns [84]. Analyzing genetic variations within mosquito populations and arbovirus populations is essential to identify high-risk areas for effective vector control measures and disease outbreak prevention [86].

Discussion

Our study underscores the significant health impact of arboviruses, particularly dengue, chikungunya, and Zika, in India [12, 19]. The epidemiological complexity of these viruses, influenced by various factors, such as immune selection and cross-reactive antibodies, poses challenges to public health in terms of diagnosis and treatment [23]. The distribution of dengue cases highlights the ongoing threat in states like Kerala, Karnataka, and Maharashtra, emphasizing the need for targeted interventions [12, 19]. The findings also emphasize the global and regional significance of vertical transmission in maintaining and spreading dengue viruses [35, 36, 38-41, 90-93]. Vector surveillance initiatives contribute to a comprehensive understanding of ZIKV and DENV prevalence in Aedes mosquitoes, crucial for effective epidemiological control [47, 48]. Furthermore, the diverse breeding habits of Aedes mosquitoes in India highlight their adaptability to varied environments [59, 60, 67].

Larval indices, such as house index, container index, and breteau index, play a crucial role in assessing dengue epidemiological risk, emphasizing the need for targeted control efforts in specific high-risk areas [60, 94].

Major ecological factors like: Climate change, global trade, travel, urbanization, and population growth have created conducive conditions for dengue vectors and viruses to proliferate [6]. Furthermore, another important factor we cannot ignore for our future vector control measures is insecticide resistance status, though an excellent review was [95] reported in India. The timeline, spanning from the dichloro diphenyltrichloroethane era to the current use of synthetic pyrethroids, reveals a concerning trend of increasing resistance patterns in India [96]. While organophosphates have shown relative efficacy, carbamates, especially temephos, crucial for larval control, exhibit alarming resistance [96]. Spatially, insecticide resistance reports are sporadic, with Delhi and Karnataka recording the highest occurrences [95]. However, comprehensive nationwide surveys are lacking, hindering a holistic understanding and highlighting the urgent necessity for an insecticide resistance management strategy, use of biocontrol agents like copepods, entomopathogenic fungi, and genetically modified mosquitoes show promise, but further research on sustainability and cost-effectiveness is crucial [97]. On the other hand, because of the knowledge gaps highlighted in Table 1 in this review, we signpost future research questions and their probable solutions that will be essential in planning surveillance and control (Figure 4) of Aedes borne diseases in India going forward:

• In-depth epidemiological studies are crucial to identify specific factors contributing to consistently high dengue cases in states [26];

• Collaborations between researchers, healthcare agencies, and vaccine developers, supported by ongoing genomic surveillance, can aid in developing a vaccine that considers the diverse antigen and genotype variations of the arbovirus [26, 28];

• Future strategies should encourage and support research initiatives in regions with limited studies, conducting ecological studies to identify factors influencing arboviral transmission variability and tailoring control measures, accordingly;

• Utilize novel molecular surveillance techniques to understand the genetic and molecular aspects of arbovirus transmission [95]; • Conduct region-specific studies to identify the temporal feeding patterns of *Aedes* mosquitoes and develop targeted control measures based on these patterns;

Conduct comprehensive studies on the toxicity of sugar baits against non-target organisms and evaluate their residual efficacy in diverse field conditions in India;

• Conduct ecological studies to understand the specific ecological and environmental factors influencing the adaptability of *Aedes* mosquitoes to diverse breeding habitats in different regions of India;

• Focus on understanding the mechanisms and frequency of mosquito movement through transportation networks, migration routes, and factors supporting longdistance dispersal [95];

• Conduct comprehensive analyses of environmental factors, such as breeding sites, land use patterns, and urbanization levels, to provide valuable insights and explain unexplained variations in genetic structure [95];

• Regular monitoring of insecticide resistance status, resistance mechanisms, and investigation of the impact of organic and other anthropogenic pollutants on the response of *Ae. aegypti* and *Ae. albopictus* to insecticides;

• Implementation of awareness initiatives like poster campaigns, volunteer education on source reduction in mosquito breeding, social media campaigns, program implementation for waste management, and essential oil-based mosquito repellent.

Conclusion

This thorough review consolidates information on the behavior, genetics, and virus transmission patterns of mosquitoes, specifically Ae. aegypti and Ae. albopictus, in India. The focus lies on crucial aspects of controlling these vectors. The research has mainly centered on their breeding habits, emphasizing the importance of understanding their feeding patterns, and host preferences for effective disease control. An innovative strategy, using attractive toxic sugar baits, shows promise in controlling mosquitoes by targeting both male and female sugarfeeding behavior. Examining arbovirus transmission dynamics, especially vertical and transovarial transmission in Aedes mosquitoes, reveals regional variations, emphasizing the need for extensive vector surveillance. Genetic diversity and population structure are crucial for assessing adaptability and disease transmission potential. Despite fewer genetic studies in India compared to

global efforts, the research underscores significant genetic diversity in *Aedes* populations, requiring continuous monitoring.

Following the guidelines from the NCVBDC, a comprehensive approach to controlling Aedes mosquitoes are essential. Hospitals should implement prevention guidelines, focusing on source reduction and environmental manipulation. Draining stagnant water from potential breeding sites is crucial. Personal protection measures, including regular use of mosquito nets and repellents, are emphasized. Eco-friendly and cost-effective larvicides, insect growth regulators, and Bacillus thuringiensis israelensis can target larvae, while indoor space spray/ fogging is effective during outbreaks. Nodal officers should monitor and coordinate efforts, ensuring effective surveillance and training. Public awareness campaigns further contribute to Aedes mosquito control. Integrating comprehensive vector surveillance with molecular investigations will inform region-specific control strategies, enhancing the precision of interventions and mitigating the impact of mosquito-borne diseases in India.

Ethical Considerations

Compliance with ethical guidelines

There were no ethical considerations to be considered in this research.

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Authors' contributions

Study design and data extraction: Preeti Acharya; Study conduction: Preeti Acharya; Drafting the manuscript: Preeti Acharya; Review, editing and final approval: All authors.

Conflict of interest

The authors declared no conflict of interest.

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