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Automated detection of ADHD: current trends and future perspective

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Abstract

Attention deficit hyperactivity disorder (ADHD) is a heterogenous pediatric disorder that affects the neurodevelopment of the frontal cortex. ADHD patients exhibit combinations of inattention, impulsiveness, and/or hyperactivity. With early treatment and diagnosis, there is potential to modify neuronal connections and improve symptoms. However, the heterogeneous nature of ADHD, combined with its comorbidities and a global shortage of diagnostic clinicians, means diagnosis for ADHD is often delayed. Hence, it is important to consider other pathways to improve the efficiency of early diagnosis, including the role of artificial intelligence. In this study, we reviewed the current literature on machine learning and deep learning studies on ADHD diagnosis and identified the various diagnostic tools used. Subsequently, we categorized these studies according to their diagnostic tool as: brain magnetic resonance imaging (MRI), physiological signals, questionnaires, game simulator and performance test, and motion data. We identified research gaps include the paucity of publicly available database for all modalities in ADHD assessment other than MRI, as well as a lack of focus on using data from wearable devices for ADHD diagnosis, such as ECG, PPG, and motion data. We hope that this review will inspire future work to create more publicly available datasets and conduct research for other modes of ADHD diagnosis and monitoring. Ultimately, we hope that artificial intelligence can be extended

to multiple ADHD diagnostic tools, allowing for the development of a powerful clinical decision support pathway that can be used both in and out of the hospital.

Keywords Attention deficit hyperactivity disorder (ADHD) · Deep learning · Machine learning · PRISMA · MRI · EEG · ECG · HRV · Questionnaires · CPT · RST · Accelerometer · Acti graphy · Pupillometric · Genetic · Social media · Artificial intelligence

1. Introduction

Attention Deficit Hyperactivity Disorder (ADHD) is a common childhood-onset neurodevelopmental condition. According to a 2016 World Health Organization-World Mental Health Surveys for 10 countries, the global prevalence rate of adult ADHD was found to be 2.8%, with a higher proportion in high-income countries, and a significant association with low education and male gender [1]. Children and adults with ADHD frequently exhibit three key symptoms: inattention, impulsivity, and hyperactivity, although symptoms are heterogeneous and individuals may display more or less of these individual symptoms, for example being classified as having inattention, or hyperactivity/ impulsivity or combined subtypes of ADHD [2].

There is increasing evidence that there are distinct differences in the structure and function of the brain in individuals clinically diagnosed with ADHD: in particular changes in neuronal connections between the specific brain regions, often accompanied by changes in brain volume on neuroimaging [3], [4] (Figure 1). These neuroanatomical differences have been linked to changes in individual's cognitive function, regulation of motivation and attention [5]. The brain's reward system, which predominantly uses the neurotransmitter dopamine, is altered in individuals with ADHD [6]. For example, the prefrontal cortex of an ADHD patient, in particular, was discovered to have abnormally low presynaptic dopamine storage [7], [8]; critically impairing the individual's attention function, cognitive process, and working memory. [7], [9].

This reward deficit syndrome has been linked to individuals with a diagnosis of ADHD being more prone to engage in behaviors that promote the production of dopamine in the brain, such as alcoholism, drug addiction, and even aggressive conduct [2]. Individuals with ADHD are, for example, twice as likely to have Substance Use Disorder (SUD) than those without [10]–[13]. For those individuals with comorbid conduct disorder, the risk of SUD is even higher - four times the rate in the general population [10].

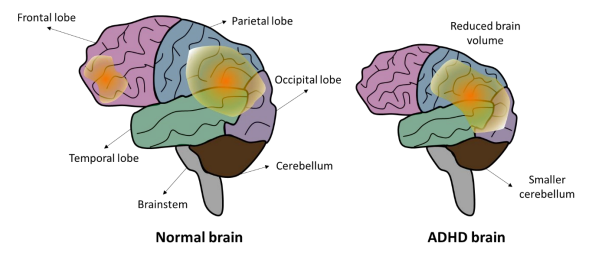


Figure 1. Schematic drawing of brains in neurotypical individuals and those diagnosed with ADHD. Yellow glowing regions represent regions of brain functional connectivity; individuals with ADHD have less neuronal connections to the prefrontal cortex as compared to normal brain.

Encouragingly, there is growing evidence that the neuroanatomical and functional changes may not be static. With appropriate early identification and treatment of ADHD symptoms, the neuroanatomy and function may resemble neurotypical individuals. For example, Mattfeld et al. [3] found that adults who had previously recovered from ADHD had restored normal brain connectivity while their minds are at rest, compared to adults who remained symptomatic with ADHD (Figure 1). Successful ADHD management can result in a significant improvement in quality of life and improved societal integration. Therefore, it is critical that ADHD is identified as early as possible, and management is evidence based, to optimize long term outcomes [14].

Currently, the diagnosis of ADHD is primarily a clinical one. An expert in ADHD diagnosis, typically a psychiatrist or specialist pediatricians, will conduct a series of clinical assessments to determine if an individual has five or more symptoms of inattention or impulsivity/hyperactivity and fulfil the DSM-5 diagnostic criteria [15]. However, clinical assessment by specialists takes a minimum of an hour, and there is a global shortages of trained specialist, meaning that diagnoses after often delayed [16]. For instance, Whitney et al. [17] reported that in Michigan, USA, there are only 11 trained psychiatrists to attend to over 100,000 children with likely mental health diagnoses. The ratio of psychiatrist-to-population is 11:100,000 for the United Kingdom, and 14:100,000 for Australia [18].

There is also evidence that adjunctive data may be helpful in diagnosing the full spectrum of individuals with ADHD, who may be overlooked or underrecognized by current clinical assessments [15]. For example, numerous studies had attempted to diagnose ADHD via neuroimaging modalities like Magnetic Resonance Imaging (MRI) [19], [20], physiological signals like electroencephalogram (EEG) [21], [22] and electrocardiogram (ECG) [23], and other modalities like accelerometers [24] and game simulators [25]. These studies aim to reduce the

workload of clinical diagnosticians by proposing artificial intelligence (AI) techniques, namely machine learning (ML) and deep learning (DL), for faster and more cost-effective ADHD diagnoses.

In this review, we aim to uncover all the different types of modalities that have been adopted by previous studies on automated ADHD diagnosis using ML or DL techniques. Machine learning is not a fully automated technique as feature extraction of the input information (e.g. MRI images, EEG, ECG, etc.) must be carried out manually, followed by feature selection of the most significant features which will ultimately be used to train the ML classifiers for automated diagnosis of ADHD [26], [27]. The DL model, on the other hand, is a fully automated process where input information can be analyzed in its original format. Hence, feature extraction and selection procedures are not mandatory in DL models [27].

2. Methods

The PRISMA guideline 2020 [28] was used in this systematic review to analyze the most relevant studies on ADHD diagnosis using either the ML or DL approach. Using the following Boolean search strings as shown in Table 1, all publications were systematically searched through PubMed, Google Scholar, IEEE, and Science Direct. All publications up to December 2021, were included in the first identification phase of the PRISMA flowchart, as illustrated in Figure 2. As a result, we began with 467 publications which was reduced to 298 after removing 165 publications with duplicated titles. Subsequently, we screened the title and abstract of the publications, and removed 165 articles that were either animal studies, conference papers, non-AI studies, non-English articles, books, review papers, and non-journal articles. We were left with 133 articles, which were downloaded and read thoroughly to assess its eligibility for this review study. Upon detailed screening of the article, we further removed more conference papers, non-journal articles, non-AI studies, review papers and irrelevant articles. We also removed articles that did not provide model accuracy results and articles which we had no access to. Finally, 91 journal articles were found eligible for inclusion in this review.

	Boolean search string					
Database	[Title]	AND [Title/Abstract]				
PubMed Google Scholar	"ADHD" OR "attention	"Madring harming" OD # lase				
IEEE	deficit hyperactivity disorder"	"Machine learning" OR "deep learning" OR "artificial intelligence				
Science direct						

Table 1. Boolean search string used for all journal article databases.

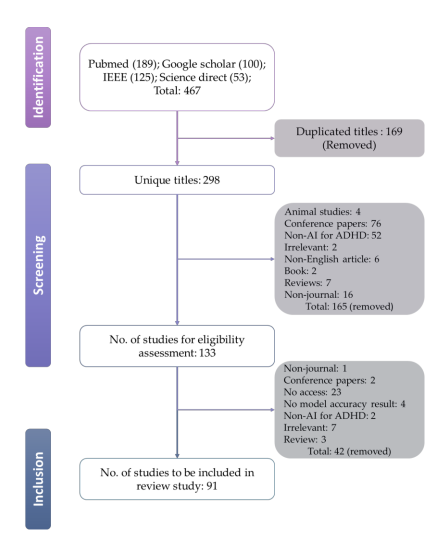


Figure 2. PRISMA flow diagram for systematic filtering of articles.

3. Results

In total, there were seven types of ADHD diagnostic tools utilized to develop AI models (Figure 3). These are discussed in the following results sections: MRI in subsection 3.1, physiological signals in subsection 3.2, questionnaire data in subsection 3.3, game simulation and performance tests in subsection 3.4, motion data in subsection 3.5, and all other studies in subsection 3.6.

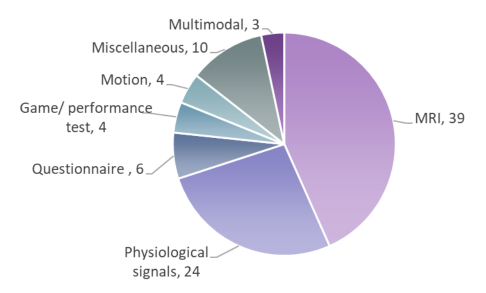


Figure 3. Pie chart representation of the ADHD assessment tools used in AI studies.

3.1 MRI

Brain MRI is the most widely studied modality for automated ADHD diagnosis, with 39 out of the 91 studies analyzing brain MRI images of ADHD patients and normal control (Table A.1). Most of the studies obtained their MRI images from one public database: the Neuro Bureau ADHD-200 Preprocessed repository (ADHD-200) [29] (Figure 4). ADHD-200 is a consortium that had collected structural and resting-state functional MRI images from 585 controls and 362 ADHD children and adolescents. Eight international imaging sites were involved in the data collection of ADHD-200, however, two out of the eight sites only provided MRI images of controls and not the ADHD individuals (Table 2). Hence, imaging data from these two sites are usually excluded from the studies. In this review, a total of 32 out of 39 MRI studies had used MRI images from ADHD-200 (Figure 4).

Table 2. Summary of number of subjects across different study sites in ADHD-200 database.

Imaging site	ADHD	Controls
Kennedy Krieger Institute	25	69
NeuroIMAGE sample	36	37
New York University Child Study Center	151	111
Oregon Health Sciences University	43	70
Peking University	102	143
University of Pittsburgh	4	94
Bradley Hospital/ Brown University	-	26
Washington University at Saint Louis	-	61
Total	361	611

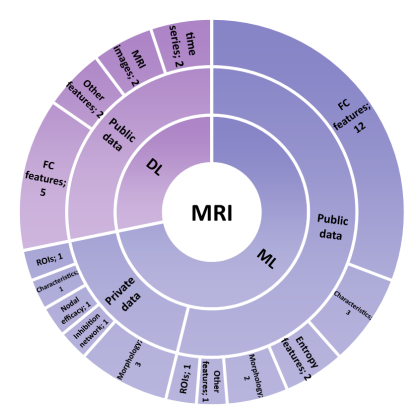


Figure 4. Sunburst plot of AI studies using MRI data. First level indicates type of AI studies, second level indicates type of dataset used, and third level indicates type of features used to train the AI models.

It is also evident in Figure 4 that ML occupied a bigger proportion than DL in the MRI analysis for ADHD; where 28 out of 39 studies had implemented ML techniques. In addition, brain functional connectivity is the most common input feature for ADHD diagnosis; 12 ML studies and 5 DL studies had utilized function connectivity (FC) features for their studies (Figure 4, Table A.1). Functional connectivity of the brain is presented in the form on a matrix, illustrating the connection between different areas of the brain [30]. Pearson correlation coefficient is commonly employed to measure if there is a strong correlation between the different brain regions, hence, resulting in a heatmap where strong and weak FC between the brain regions is evident [30].

3.2 Physiological signals

Twenty-four studies utilized physiological signals to detect ADHD, most commonly electroencephalogram (EEG: 23 studies) and electrocardiogram (ECG: 1 study) (Figure 5, Table A.2). We also observed that studies using physiological signals for the detection of ADHD had high model performances; all models had accuracy results above 80% for ML and DL (Table A.2). Only one [31] out of the 24 studies had used a public EEG database: the National Brain Mapping Laboratory of Iran [32]. The rest had used their own private datasets.

As for the type of feature most extracted from EEG signals, seven ML and three DL studies had attempted to obtain power spectral features (Figure 5). Spectral analysis of EEG involves decomposing the signal into medically established frequency sub-bands, namely, alpha rhythm (8–13 Hz), beta rhythm (13–30 Hz), delta rhythm (1–4 Hz), theta rhythm (4–8 Hz), and gamma rhythm (30–80 Hz) [33]. These frequency sub-bands are evidently different between children with ADHD and controls. A study by Kamida et al. [34] discovered that children with ADHD have higher beta activity in all brain regions except for the occipital region. Another study [35] which investigated the power spectral differences between ADHD of the inattentive type and the combined type found higher theta and alpha activities in the combined type, while higher theta/beta ratio was observed in the inattentive type.

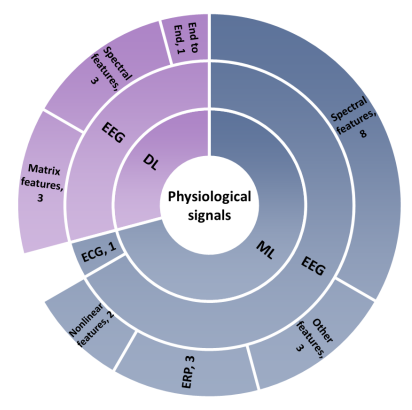


Figure 5. Sunburst plot of AI studies using physiological signals. First level indicates type of AI studies, second level indicates type of physiological signal, and third level indicates type of feature used to train the AI models.

There is only one study which utilized ECG signals (Figure 5). Even though ECG does not provide direct information on the brain activity, the autonomic nervous system links the brain to body interaction, causing fluctuations in physiological signals like ECG when an individual senses danger. For instance, an individual under acute stress will have significant increase in the heart rate (ECG) and the same phenomenon was also observed in ADHD individuals [23], [36]. Koh et al. [23] proposed ensemble ML classifier with entropy features extracted from ECG signals and detected ADHD individuals with high classification accuracy of 87.2%.

3.3 Questionnaires/ rating scales

There are various types of questionnaires or rating scales that medical professionals use to diagnose ADHD. In this section, there are only six studies that analyzed questionnaire/ rating scales data and only ML models were proposed (Table 3). It can also be seen in table 3 that decision tree (DT) classifier, including random forest classifier, are commonly proposed to analyze questionnaire data. We will only cover the questionnaires that studies have utilized to develop their best performing models (Table 3).

- **Conners' Rating Scales** are widely implemented to assess the social impact of ADHD, for example an individual's behavior in school or work [37]. Conners' parent rating scales (CPRS) used by Bledsoe et al. [38] is a parentally completed report, while Conners' adult ADHD rating scales (CAARS) used by Christiansen et al. [39], is a self-reported questionnaire.
- **Diagnostic Interview for ADHD in adults (DIVA)** [40] adopted by Tachmazidis et al. [41], is a semi-structured interview constructed based on Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition (DSM-IV) criteria for ADHD diagnosis. The interview aims to assess the symptom of ADHD in five aspects of daily life: social contact; hobbies; self-confidence; relationships; work and education.
- Behavior Rating Inventory of Executive Function Preschool version (BRIEF-P) is a 63item questionnaire for parents or teacher to rate the child's executive functions such as emotions control, working memory, organization and planning skills [42]. This questionnaire was utilized by Öztekin et al. [43] to develop their ML model.
- Adult ADHD Self-Report Scale (ASRS) used by Kim et al. [44] is created by the World Health Organization and it consists of 18 items based on the DSM-IV criteria. ASRS is a symptoms checklist for individuals to self-evaluate if they exhibit any symptoms relating to ADHD [45].
- Minnesota Multiphasic Personality Inventory-2 (MMPI-2) is a 567-item questionnaire where individuals are only required to answer 'true' or 'false' [46]. MMPI-2 is also used by Kim et al. [44], alongside ASRS to develop their ML model, and it is widely implemented to assess various mental health problem apart from ADHD such as depression, anxiety, and psychopathy [46].
- Social Responsiveness Scale (SRS) is a 65-item questionnaire that attempts to measure the social ability of individuals between ages 4 to 18 years [47]. This questionnaire is adopted by Duda et al. [48] to differentiate ADHD individuals from patient with Autistic Spectrum Disorder (ASD).

Author [ref]	Private datasets	Questionnaires	ML model	Accuracy
Bledsoe et al. [38]	23 ADHD	CPRS	SVM +DT	100
[]	12 normal			
Tachmazidis et al.	45 ADHD male	DIVA	DT +	95.7
[41]	24 ADHD female	DIVA	knowledge	95.7
Vim at al [44]	5726 college		Random	93.6
Kim et al. [44]	students	MMPI+ASRS	forest	93.0
Öztekin et al. [43]	87 ADHD	BRIEF-P	SVM	92.6
Oztekin et al. [45]	75 normal			92.0
Duda at al [49]	174 ADHD	SRS	ENet and	82.0
Duda et al. [48]	248 ASD	5K5	LDA	82.0
	385 ADHD			
Christiansen et al.	135 Obesity		DT	
[39]	517 problematic	CAARS	21	80.0
	gambling		(lightGBM)	
	592 normal			

Table 3. Summary of AI studies that used questionnaire data to develop AI model.

3.4 Game simulation and performance tests

This section discusses the use of conventional performance tests and game simulation to diagnose ADHD. There are two ML studies each, which utilized performance tests and game simulation respectively, to train their model (Table 4). Continuous Performance Test (CPT) and Reverse Stroop task (RST) are neuropsychological tests to evaluate the selective and sustained attention of an individual [49], [50]. The CPT is a computerized test which requires participants to react correctly to a specific stimulus [25]. For instance, participants are told to press the spacebar for all letters except for 'O'. In traditional Stroop task, participants are given words, for example 'Blue', which can be presented in different colors: 'Blue' (incongruent color red), 'Blue' (congruent color blue), and 'Blue' (neutral color black). Participants are then required to provide the color of the word, instead of the meaning of the word. Hence, in RST, the task is reversed where participants have to read out the meaning of the word regardless of the color it is printed in [51].

As for game simulations, its main purpose is creating an interactive environment that is customizable to best suit the user's needs [25], [52]. Yeh et al. [52] created a virtual reality (VR) classroom and incorporated a series of tests, including CPT, for ADHD diagnosis. In their VR system, some 'distractions' such as 'teacher standing up', 'door open', or 'thunder shower', were also included. They then recorded the test results, reaction time, and focus time for the user to complete the test. On the other hand, Heller et al. [25] utilized a videogame known as 'Groundskeeper' that is specially developed by CogCubed [53] for early ADHD detection. They extracted 33 game data variables, and trained four different ML classifier: random forest,

AdaBoost, J48, and JRip. However, they did not specify which classifier provided the best performance result.

Author [ref]	Private	Mode	Tests	ML models	Accuracy
	datasets				
Slobodin et	213 ADHD	Performance	CPT	Random	87.0
al. [54]	245 normal			Forest	
Yasumura et	108 ADHD	Performance	RST	SVM	86.3
al. [55]	108 normal				
Yeh et al. [52]	37 ADHD	Game	VR system	SVM	83.2
	31 normal				
Heller et al.	26 ADHD	Game	Groundskeeper	-	78.0
[25]	26 normal		(CogCubed)		

Table 4. Summary of list of AI studies that used standard performance test or game simulation data to develop their model.

3.5 Motion data (actigraphy & accelerometer)

Motion activity can also be a diagnostic marker for ADHD. In this section, two types of motion activity measure - actigraphy and accelerometer - are covered along with the four studies that utilized these motion data as listed in Table 5. Both actigraphy and accelerometer data are recorded via an accelerometer device that is usually worn on the wrist of the dominant arm and ankle of the dominant leg [56], [57]. The difference between the studies that analyzed the two types of motion data is the type of activity the subject is doing; actigraphy studies the subject's sleep efficiency [56], whereas accelerometer analyzes the subject's motion during normal daily activities [57]. As such, there are some studies that reported ADHD patients exhibited more movement than the controls during sleep [58] which correlates to increased daytime sleepiness [59]. This demonstrates that increased activity level is a well-known feature of ADHD, which is also reflected in their daily routine, and can be easily monitored with wrist-worn accelerometer devices [60], [61].

In either case, the accelerometer device used to record the motion data is designed to be unobtrusive, allowing the participants to be natural in their own environment. This would not be possible with EEG or polysomnography recording procedures because data collection takes place in a laboratory that the participants are unfamiliar with, and they are also required to attach a large number of electrodes, which can be very uncomfortable [62]. This, in turn, may have an impact on the quality of data collected.

Table 5. Summary of AI studies that used actigraphy or accelerometer data.

Author [ref] Private datasets	Mode	Features	Models	Accuracy
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Faedda et al. [56]	44 ADHD 21 ADHD+depression 48 bipolar 42 controls	Actigraphy	28 metrics	ML (SVM)	83.1
Amado- Caballero et al. [57]	73 ADHD 75 normal	Accelerometer	end-to-end	DL (CNN)	98.6
O'Mahony et al. [24]	24 ADHD 19 normal	Accelerometer	inertial measurement units	ML (SVM)	95.1
Muñoz- Organero et al. [63]	11 ADHD 11 normal	Accelerometer	Acceleration image	DL (CNN)	93.8

3.8 Miscellaneous (pupillometric, twitter and genetic)

In this section, we cover the least common modalities of ADHD diagnosis that ML studies have used. Two ML studies had utilized pupillometric data while only one study had analyzed twitter data (Table 6).

Interestingly, studies have shown that the brain norepinephrine system which is associated to pupil-size dynamics, is found to be impaired in ADHD patients [64]. Another study has also demonstrated that ADHD patients (off-medication) have decreased pupil diameter when performing visuo-spatial working memory task as compared to the controls [65]. This could be due to the difficulty in suppressing saccadic eye movements in ADHD patients who need to fixate [66]. Hence, uncontrollable eye movement in ADHD patients can be a potential biomarker for diagnosis, as Varela Casal et al. [66] and Das et al. [64] have implemented in their ML studies.

With the rise of social media, Twitter has become a potential platform of ADHD detection among the Twitter users [67]. A majority of the mentally ill are reluctant to seek help from mental health care professionals, which results in gradual accumulation of suicidal thoughts in the absence of professional help [68]. Hence, social media platform like Twitter, has become a source of comfort for these individuals to discuss mental health issues openly, as they seek connection and support from people of the same community [69]. Therefore, social media platform can be utilized for early detection of various mental illnesses and intervene suicidal actions [69]. In the study by Guntuku et al. [70], they identified highly correlated topics in Twitter and used it as a learning feature for their support vector machine (SVM) classifier (with 76% accuracy).

Table 6. Summary of AI studies that used pupillometric or Twitter data to develop their model.

Author [ref]	Private	Modality	Feature	ML model	Accuracy
	datasets				

Varela Casal	21 ADHD	Pupillometric	Eye Vergence	SVM	96.3
et al. [66]	21 normal				
Das et al. [64]	28 ADHD	Pupillometric	pupil-size	SVM	76.1
	22 normal	_	dilation		
			velocity and		
			acceleration		
			feature		
Guntuku et	1032 ADHD	Twitter	Topic	SVM	76.0
al. [70]	1029 normal				

There is a known genetic influence on the likelihood that an individual will be diagnosed with ADHD [71], [72]. Numerous twin studies have reported a high heritability estimate of approximately 80% for both monozygotic and dizygotic twins [73]. ML and DL have recently been applied in seven studies to help identify ADHD genetic biomarkers. A summary on the four ML and three DL studies for ADHD are listed in Table 7. However, it is important to note that the genetic biomarkers identified [74]–[78] in Table 7 did not follow the standard genome-wide association studies (GWAS), which identify risk genetic variants via its significant P-values (i.e. not be lower than 5×10^{-8}) [75].

Author [ref]	Dataset	Model	Findings
Sokolova et al. [74]	87 ADHD	ML (Bayesian	DAT1 risk haplotype
	77 normal	Constraint-based	only has direct
		Causal Discovery	influence on the
		algorithm)	ADHD inattentive
			type.
Liu et al. [75]	1033 ADHD	DL (CNN)	EPHA5 is identified
	950 normal		as a potential risk
			gene of ADHD.
			Model diagnostic
			accuracy = 90.2%
Liu et al. [76]	116 ADHD	DL (MLP)	GRM1 and GRM8
	408 normal		genes are identified
			to have the highest
			weight in ADHD
			diagnosis
			Model diagnostic
			accuracy = 78.0%

Table 7. Summary of AI studies that used genetic data to develop their model and identify ADHD genetic variant.

Esteller-Cucala et al.	20,000 ADHD	DL (Approximate	Frequency of ADHD-
[79]	35,000 normal	Bayesian	risk alleles decreased
		Computation	since ancient time
		coupled + deep	and have become
		learning framework)	maladaptive in
			today's society.
Cervantes-Henríquez	408 ADHD	ML (ensemble)	The proposed model
et al. [77]			identifies ADGRL3,
			DRD4, and SNAP25
			genes as contributing
			to the severity of
			ADHD.
Sudre et al. [80]	362 ADHD	ML (Random Forest)	Participants with the
			highest polygenic
			risk score for ADHD
			usually have
			worsening
			symptoms.
Jung et al. [78]	39 ADHD	ML (SVM)	The proposed model
	34 normal		identified the COMT
			gene as having an
			impact on the
			abnormal
			development of the
			frontal cortex in
			ADHD patients.

4. Discussion

The 'gold standard' to diagnose ADHD usually relies on a combination of neuropsychological tests, rating scales, behavioral observations, examinations, and evaluation of the impact of treatment trials [81]. This is time consuming and limited by the number of trained diagnostic specialists globally. We reviewed the accuracy of the application of ML and DL to a range of well-established diagnostic tools, such as questionnaires/ rating scales, as well as more innovative diagnostic tools, including MRI and EEG, as summarized in Table 8. All but three ML studies adopted single modality approaches for ADHD diagnosis, however a multimodal approach may be well suited to ADHD, due to its heterogeneous clinical nature.

Table 8. Summary of AI studies using multiple modalities to develop their model.

Author	Private	Modality	Feature	ML model	Accuracy
[ref]	datasets				

Yoo et al.	191 ADHD	fMRI+genetic	cortical thickness	RF	85.1
[82]	78 normal		and volume		
			features		
Kautzky et	16 ADHD	Genes+PET+MRI	SNPS+ROI	RF	82.0
al. [83]	22 normal				
Crippa et	22 ADHD	blood+EEG+	neuropsychological,	SVM	81.0
al. [84]	22 normal	cognitive test	FA profiles, and		
			deoxygenated-		
			hemoglobin		
			features		

It is estimated that 60 to 100% of ADHD children will develop one or more comorbid mental health or behavioral disorders as they reach adulthood [85], [86] including conduct disorder, depression, autism spectrum disorder (ASD) and bipolar disorder. The presence of comorbidities can make accurate diagnosis even more challenging [87]. An accurate diagnosis is required in order to tailor appropriate therapies. The preliminary evidence reviewed in this study suggests that AI can play a helpful role in diagnosing individuals with ADHD with and without comorbidities. There are nine ML studies in this review that had attempted to differentiate ADHD from other mental disorders or diagnose ADHD in individuals with a range of comorbidities (Table 9).

Author [ref]	Dataset	Modality	Comorbid condition	ML Model	Accuracy (%)
Tor et al. [88]	45 ADHD 62 ADHD+CD 16 CD	EEG	Conduct disorder	kNN	97.88
Vaidya et al. [89]	307 ADHD 240 ASD 465 Control	MRI	ASD	SVM	88.9
Koh et al. [23]	45 ADHD 62 ADHD+CD 16 CD	ECG	Conduct disorder	bagged tree classifier	87.2
Jun et al. [90]	86 ASD 83 ADHD 125 normal	MRI	ASD	SVM	84.1
Faedda et al. [56]	44 ADHD 21 ADHD+depression 48 bipolar 42 controls	Actigraphy	Bipolar depression	SVM	83.1
Duda et al. [48]	174 ADHD 248 ASD	Questionnaire	ASD	ENet and LDA	82.0

Table 9. Summary of AI studies that had considered other comorbidities of ADHD.

Christiansen et al. [39]	385 ADHD 135 Obesity 517 problematic gambling 592 normal	Questionnaire	Obesity, problematic gambling	DT (lightGBM)	80.0
Heller et al. [25]	26 ADHD 26 normal	Game	Depression, ASD, anxiety, disruptive behavior disorder	-	78.0
Guntuku et al. [70]	1032 ADHD 1029 normal	Twitter	Depression Bipolar Anxiety	SVM	76.0

In total, we report on 68 ML studies and 23 DL studies for ADHD diagnosis. SVM is the most commonly used classifier in ML research, while convolutional neural network (CNN) is the most commonly proposed model in DL research (Figure 6). This is not to say that SVM or CNN are superior to other ML or DL models. The suitability of an ML or DL model is determined by the type of dataset and feature used to train the classifier, while the practicality of the ML or DL model is determined by a well-structured clinical trial in which their models are tested in real clinical settings with direct interaction with ADHD patients [27]. There is definitely scope for improvement with AI methodology for ADHD diagnosis before it can be considered for clinical use. From Figure 7, we can see that DL research started only in 2017 and has yet to reach maturity in its technological advancement, whereas the percentage of ML studies has declined with the rise of DL research. This is not surprising given the large number of ML studies in ADHD diagnosis, which has made it extremely competitive and difficult for new studies to outperform previous ones. The average model accuracy reported by these ML and DL studies has remained stable between 80 and 90% since 2013. Nonetheless, the rapid increase in AI studies in recent years, as technology advances, indicates that computer-aided ADHD diagnosis is improving. As such, we hope to encourage more DL studies in ADHD diagnosis so that its feasibility can be demonstrated, and a clinical trial can be conducted.

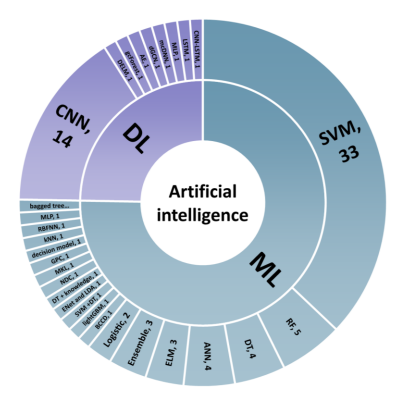


Figure 6. Sunburst plot of AI studies analyzed in this review. First level indicates type of AI studies, and second level indicates type of classifier proposed by AI studies.

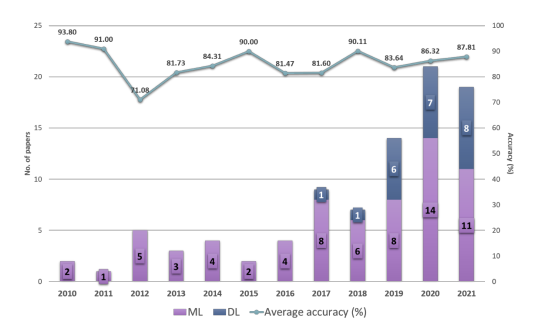


Figure 7. Bar chart representation of the number of AI studies in ADHD diagnosis published across the years from 2010 to 2021. Line graph represents the average model accuracy of AI studies across the years.

In summary, the significance of this review is as follows:

- We have followed PRISMA guideline and gathered 91 AI studies for this review.
- From the 91 studies, we have categorized them according to the type of modality or dataset used to train their model for ADHD diagnosis. Namely, MRI, physiological signals, questionnaires, game simulation, performance test, motion data, and miscellaneous which includes pupillometric, twitter and genetic data.
- For modalities like MRI and EEG signals, which are adopted by a large number of studies in this review, we have identified the most commonly used features for ADHD diagnosis, which is functional connectivity features and power spectral features for MRI and EEG respectively.
- For studies that have used questionnaire/ rating scales data, we have listed the standard ADHD assessment questionnaires that have successfully helped ML models to achieve high diagnostic performance.
- We have also identified ML studies that have utilized parameters from computerized neuropsychological tests like RST and CPT for ADHD diagnosis. This could reduce the burden on clinical diagnosticians if AI can analyze the performance result in their place.
- Two studies had attempted to create a customizable environment for ADHD diagnosis. One study had also created a virtual reality environment to conduct a CPT test [52]. Another study had used game parameters for ADHD diagnosis [25].
- We identified a small number of studies that used motion data from accelerometers and actigraphy- they prioritized subjects' comfort and collected motion data in the subjects' natural environment.
- Lastly, we identified a small number of studies using pupillometric and Twitter data. We also found a few studies that used both ML and DL on genomic sequencing data to help identify genetic biomarkers for ADHD.

This review also has some limitations:

- Apart from the popular ADHD-200 MRI database, the scarcity of large publicly available ADHD databases for the rest of the modality category caused majority of the studies in this review to use private datasets.
- For private datasets, the number of subjects and the methods used to collect data varied greatly.
- Even for study that had used ADHD-200 database, the number of subjects included in their studies varied greatly as well.
- It is difficult to compare the results of ADHD studies as different datasets were used, and there was a great variation in the number of subject and data acquisition methods.

5. Future direction for AI in ADHD diagnosis.

There are several future research pathways that could be followed to further explore the use of AI as a clinical decision support tool for ADHD. One optimal pathway could be the use of a cloud system, as depicted in Figure 8, that has unified data covering all ADHD diagnostic tools. As a result, clinicians could have ready access to all the information needed to confirm a diagnosis. For example, parents, teachers, or ADHD patients could complete the questionnaires on their own and have the AI models in the cloud system analyze the questionnaire data for psychiatrists or pediatricians. Questionnaires have the potential to be open to bias, and somewhat subjective interpretation, whereas wearable devices, on the other hand, may be able to provide objective measurements that provide a useful adjunct to ADHD diagnosis. There is also scope for researchers to apply AI not only as part of a diagnostic pathway, but also as a precision medicine clinical decision support system to help with tailoring and monitoring treatments.

Firstly, it is critical that more publicly accessible ADHD databases are available. Other than the publicly available MRI ADHD-200 database, which was used by 32 studies in this review, other studies needed to utilize private datasets. Since ADHD is a heterogenous disorder, it is important that databases of different types of ADHD diagnostic tools are available. In particular physiological signals (ECG) and motion data (accelerometer), would be highly desirable, as these were the biological tools that have shown more effectiveness representing the inattentiveness, impulsiveness, and hyperactivity symptoms of individuals during their daily routines.

Secondly, further exploration of the utility and effectiveness of wearable devices for ADHD diagnosis and monitoring would be warranted. Currently, only ECG signals and motion signals gathered from accelerometers are suitable to be collected from wearable devices. However, very few AI studies have attempted to use these data to diagnose ADHD; only one study used ECG signals, while four studies used motion data. There is also another study that had used heart rate variability (HRV) for ADHD diagnosis, but no accuracy data was reported hence this study was not included in this review [91]. Nonetheless, this shows that HRV is also another possible parameter for ADHD diagnosis. Another parameter that could be explored is photoplethysmography (PPG) signals which can easily be acquired from smartwatches, smartphones and oximeters [92]. An advantage of PPG signals is that they have low bandwidth requirements which do not deplete battery's capacity excessively [93], making them excellent candidates for signals storage in a cloud system, as shown in Figure 8. Ambulatory signal collection is also very helpful in telehealth situations, which is increasingly being used clinically since the COVID-19 pandemic.

Furthermore, despite the large amount of AI research in MRI and EEG for ADHD diagnosis found in this review, it is important to note that neither diagnostic tool is currently used by psychiatrists or pediatricians to diagnose ADHD in routine clinical settings [94], [95]. As both diagnostic tools can only be used in hospital settings or laboratories, gathering data from individuals with a diagnosis of ADHD is a time-consuming and expensive process [95], [96]. MRI requires patients to be still, which can be particularly challenging for people with ADHD, resulting in a higher chance of motion artifacts that render interpretation very difficult [97], [98]. Therefore, wearable devices such as PPGs or accelerometers may be better suited as data acquisition devices for ADHD patients and should be investigated further for future AI studies.

AI methodologies like ML and DL can suffer from poor interpretability [99], [100]. Because of the complicated algorithm used to derive the result, DL models are referred to as a "black box", and clinicians find it hard to understand their outputs. The poor interpretability of AI algorithms has hampered their adoption in healthcare as a clinical decision support tool [101]. Therefore, future work for DL models should focus on the explainability of the model. For instance, there are a few techniques such as LIME, SHAP or integrated gradients, that can improve the interpretability of ML or DL models [102]. We hope to encourage the development of a practical AI model for ADHD diagnosis and monitoring, which will be an important component of the cloud system depicted in Figure 8.

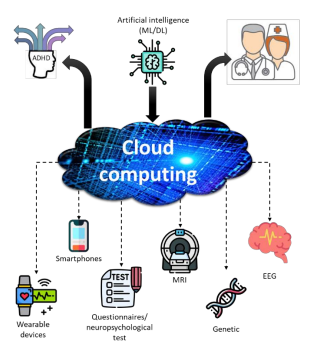


Figure 8. Cloud system designed for ADHD diagnosis and monitoring.

6. Conclusion

This review surveyed various ADHD diagnostic tools and included studies that used ML and DL AI techniques to perform the diagnosis. Ninety-one studies were determined to be eligible for this review, and they were further subdivided into their respective modalities for critical analysis. As a result, we notice that the majority of the studies were inclined towards hospital settings modalities like MRI and EEG, while the rest of the modalities were reported by very few studies. In addition, there was lack of publicly available dataset for the majority of the modalities except for MRI. There were limited studies using data acquired from wearable devices like ECG and accelerometer, and there were no studies that attempted to use PPG signals. Therefore, we propose that future research should focus on developing more publicly available datasets for the

other modalities in ADHD assessment and develop AI models that utilized data from wearable devices for ADHD diagnosis and monitoring. We also suggest future AI studies in ADHD to improve the interpretability of their models to encourage adoption in healthcare. With more robust research in AI techniques, a cloud system capable of reaching out to various ADHD diagnostic tools could become a reality and serve as an indispensable clinical decision support tool for clinicians.

Declaration of Competing Interest

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

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Author	Year	Dataset	Subjects	Feature extracted	Classifier	Accuracy (%)			
	Deep learning (DL)								
Zhang et al. [103]	2020	public (Neuro Bureau ADHD-200 dataset)	422 ADHD 597 normal	time-series signals	CNN	54.1			
Zou et al. [104], 2017	2017	public (Neuro Bureau ADHD-200 dataset)	285 ADHD 491 normal	functional connectivity + morphology feature	CNN	69.2			
Mao et al. [105], 2019	2019	public (Neuro Bureau ADHD-200 dataset)	359 ADHD 429 normal	preprocessed fMRI scans	CNN	71.3			
Zhao et al. [106]	2021	public (Neuro Bureau ADHD-200 dataset)	260 ADHD 343 normal	functional connectivity features	dGCN	72.0			
Peng et al. [107]	2021	public (Neuro Bureau ADHD-200 dataset)	351 ADHD 430 normal	functional connectivity features	CNN	72.9			

Appendix table A.1. Summary of list of AI studies that used MRI data to develop their model.

					1	
Riaz et al. [108]	2020	public (Neuro Bureau ADHD-200 dataset)	351 ADHD 430 normal	fMRI time-series signals	CNN	73.1
Chen et al. [109]	2019	public (Neuro Bureau ADHD-200 dataset)	362 ADHD children 585 normal children	combination of imaging and personal characteristic data	mcDNN	78.3
Shao et al. [110]	2019	public (Neuro Bureau ADHD-200 dataset)	310 ADHD 359 normal	functional connectivity features	gcForest	82.7
Khullar et al. [111]	2021	public (Neuro Bureau ADHD-200 dataset)	351 ADHD 430 normal	Raw images	CNN- LSTM	98.2
Preetha et al. [112]	2021	public (Neuro Bureau ADHD-200 dataset)	260 ADHD- C children 173 ADHD- I children 744 normal children	-	DELM	98.2
Tang et al. [20]	2021	public (Neuro Bureau ADHD-200 dataset)	-	functional connectivity features	AE	99.6
			Machine lea	arning (ML)		
Colby et al. [113]	2012	public (Neuro Bureau ADHD-200 dataset)	285 ADHD 491 normal	functional connectivity features, Structural and morphological features	SVM-RBF	55.0
Qureshi et al. [114]	2016	public (Neuro Bureau ADHD-200 dataset)	67 ADHD- C children 67 ADHD-I children 67 normal children	Cortical Thickness and volume features	ELM	60.8
Brown et al. [115]	2012	public (Neuro Bureau ADHD-200 dataset)	192 ADHD- C children 124 ADHD- I children 523 normal children	characteristic data	Logistic	62.5
Zhou et al. [116]	2021	Private	116 ADHD 116 normal	macrostructural property, Morphometric measures, Image intensity measures	MKL	64.3

[1]
Itani et al. [117]	2019	public (Neuro Bureau ADHD-200 dataset)	146 ADHD 105 normal	gender and 26 ROI	DT	66.6
Anderson et al. [118]	2014	public (Neuro Bureau ADHD-200 dataset)	276 ADHD 472 normal	Phenotypic, Independent Components, motion, structural, functional connectivity features	DT	66.8
Sato et al. [119]	2012	public (Neuro Bureau ADHD-200 dataset)	249 ADHD 122 ADHD- I	functional connectivity features	Logistic	67.0
Tan et al. [120]	2017	public (Neuro Bureau ADHD-200 dataset)	117 ADHD 98 normal	FV and demographic variables	SVM	68.6
Sidhu et al. [121]	2012	public (Neuro Bureau ADHD-200 dataset)	141 ADHD- C children 98 ADHD-I children 429 normal children	phenotypic + imaging data	SVM	72.9
Chaim- Avancini et al. [122]	2017	Private	52 ADHD 44 normal	ROIs	SVM	73.8
Wang et al. [123]	2018	private	36 ADHD 35 normal	interregional morphological patterns	SVM-RFE	74.6
Liu et al. [124]	2020	public (Neuro Bureau ADHD-200 dataset)	351 ADHD 430 normal	Deep learning model extracted features	AdaDT	75.6
Luo et al. [125]	2020	Private	36 ADHD 36 normal	Features of nodal efficiency	Ensemble	76.6
Hart et al. [126]	2013	Private	30 ADHD 30 normal	inhibition networks	GPC	77.0
Khan et al. [127]	2021	public (Neuro Bureau ADHD-200 dataset)	295 ADHD 364 normal	functional connectivity features	SVM	81.0
Miao et al. [128]	2019	public (Neuro Bureau ADHD-200 dataset)	308 ADHD 361 normal	Principle Components and Entropy-Based Features	DT	81.8
Jun et al. [90]	2018	public (ABIDE and ADHD200 dataset)	86 ASD 83 ADHD 125 normal	ROI-to-ROI functional connectivity feature	SVM	84.1

Sun et al. [129]	2020	public (Neuro Bureau ADHD-200	351 ADHD 430 normal	functional connectivity features	SVM	85.3
Shao et al. [130]	2020	dataset) public (Neuro Bureau ADHD-200 dataset)	35 ADHD 32 normal	Principle Components and Entropy-Based Features	T-R-SVM	86.4
Riaz et al. [131]	2017	public (Neuro Bureau ADHD-200 dataset)	59 ADHD 93 normal	functional connectivity features	SVM	86.8
Chen et al. [132]	2020	public (Neuro Bureau ADHD-200 dataset)	272 ADHD 361 normal	functional connectivity features	SVM	88.1
Vaidya et al. [89]	2019	private	307 ADHD 240 ASD 465 Control	3 behavioral profiles	SVM	88.9
Deshpande et al. [133]	2015	public (Neuro Bureau ADHD-200 dataset)	260 ADHD- C children 173 ADHD- I children 744 normal children	linear/nonlinear directional/nondirectional functional connectivity features	FCC ANN	90.0
Peng et al. [134]	2013	public (Neuro Bureau ADHD-200 dataset)	59 ADHD 93 normal	brain structure features	ELM	90.2
Qureshi et al. [135]	2017	public (Neuro Bureau ADHD-200 dataset)	67 ADHD- C children 67 ADHD-I children 67 normal children	functional connectivity features	ELM	92.9
Johnston et al. [136]	2014	private	34 ADHD 34 control	white matter images (m)	SVM	93.0
Tang et al. [137]	2020	public (Neuro Bureau ADHD-200 dataset)	59 ADHD 93 normal	functional connectivity features	Decision model	97.6
Bohland et al. [19]	2012	public (Neuro Bureau ADHD-200 dataset)	-	Gender, Non-Imaging Phenotypic, Anatomical, and functional connectivity features	SVM	98.0

Author	Year	Dataset	Subjects	Feature extracted	Sampling frequency	Classifier	Accuracy (%)	
			EEG – Dee	p learning (DL))			
Vahid et al. [138]	2019	private	48 ADHD 44 normal	end-to-end	500	CNN	83.0	
Dubreuil- Vall et al. [139]	2020	private	20 ADHD 20 normal	EEG spectograms	500	CNN	88.0	
Chen et al. [140]	2019	private	50 ADHD 57 normal	Grad-CAM	1000	CNN	90.3	
Tosun et al. [141]	2021	private	1088 ADHD sample 1088 normal sample	Power spectral features	500	LSTM	92.2	
Chen et al. [142]	2019	private	50 ADHD children 51 normal children	connectivity matrix	1000	CNN	94.7	
Moghaddari et al. [31]	2020	public (National brain mapping laboratory of Iran)	31 ADHD 30 normal	Power spectral band separation Making RGB images	128	CNN	98.5	
Ahmadi et al. [22]	2020	private	13 ADHD- C children 12 ADHD-I children 14 normal children	spatial and power spectral features	250	CNN	99.5	
	EEG – Machine learning (ML)							
Müller et al. [143]	2019	private	181 ADHD 147 normal	Power spectral features, ERP peak amplitudes and latencies	500	SVM	80.0	
Kim et al. [144]	2021	Private	34 ADHD 45 normal	MMN source activity features	1000	SVM	81.0	

Appendix table A.2. Summary of list of AI studies that used physiological signals to develop their model.

			1					
Tenev et al. [145]	2014	private	67 ADHD 50 normal	eye close, eye open, ECPT and VCPT	256	Ensemble	82.3	
Khoshnoud et al. [146]	2018	private	12 ADHD 12 normal	nonlinear power spectral features	256	SVM	83.3	
Chen et al. [147]	2019	private	50 ADHD 58 normal	Power spectral features	1000	SVM	84.6	
Mueller et al. [148]	2011	private	75 ADHD 75 normal	ERP components	250	SVM	91.0	
Altınkaynak et al. [149]	2020	private	23 ADHD 23 normal	Morphological, nonlinear, and wavelet features	2500	MLP	91.3	
Mueller et al. [150]	2010	private	74 ADHD 74 control	ERP components	-	SVM	92.0	
Ahmadlou et al. [151]	2010	private	40 ADHD 7 normal	Power spectral features	256	RBFNN	95.6	
Tor et al. [88]	2021	private	45 ADHD 62 ADHD+CD 16 CD	nonlinear features	500	kNN	97.9	
Guney et al. [152]	2021	private	27 ADHD 38 normal	event-related potentials (ERPs)	1000	ANN	98.4	
Rezaeezadeh et al. [153]	2020	private	12 ADHD children 12 normal children	non-linear power spectral features	256	SVM (RBF)	99.6	
Joy et al. [154]	2021	private	5 ADHD 5 normal	nonlinear power spectral features	256	ANN	99.8	
Kaur et al. [155]	2019	Private	47 ADHD 50 normal	PSR-PSO	256	NDC	100.0	
Bashiri et al. [156]	2018	private	15 ADHD- HI 30 ADHD-I	Power spectral features	250	ANN	100.0	
Öztoprak et al. [21]	2017	private	70 ADHD 38 normal	Power spectral features	1000	SVM-RFE	100.0	
	ECG – Machine learning (ML)							
Koh et al. [23]	2021	private	45 ADHD 62 ADHD+CD 16 CD	entropy features	500	Ensemble	87.2	

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