

author	year	design	sampling	Exclusion criteria	Inclusion criteria	Recruitment	consecutive	duration	clinical condition	Type of measurement
Bove	2015	Cohort study, paired control-group, feasibility study, human observational trial	Convenient	not reported	Informed consent, Multiple sclerosis	Pairs consisting of 1 patient with demyelinating disease and 1 healthy cohabitant, were recruited at the Partners MS Center, a large referral clinical center in the northeastern United States.	no	1 year monitoring	Severity of MS	Questionnaires and visual tests should classify severity of MS
Bove	2015	Cohort study, paired control-group, feasibility study, human observational trial	Convenient	not reported	Informed consent	Pairs consisting of 1 patient with demyelinating disease and 1 healthy cohabitant, all aged 18–55 years, were recruited at the Partners MS Center, a large referral clinical center in the northeastern United States. Cohabitant pairs were recruited to control for common environment.	no	1 year monitoring	Healthy cohabitants of participating patients	Questionnaires and visual tests should provide a control for environmental factors
Kaiser	2013	Open-label, single-arm, multicentre study, Pilot study	Prospective	Patients were excluded if they had concomitant ocular disease in the study eye; neurologic impairment that would interfere with study assessments; use of systemic medications known to be toxic to the lens, retina, or optic nerve; or use of any other investigational agents within 60 days of screening.	Active CNV secondary to AMD (either newly diagnosed and treatment-naive or successfully treated with anti-vascular endothelial growth factor therapy for 1 year) in at least 1 eye, eligible for ranibizumab therapy, with best corrected visual acuity (BCVA) letter score 24 or higher (20/320 Snellen equivalent) by "Early Treatment Diabetic Retinopathy Study chart" at 4 m. Only one eye was required to fulfill entry criteria for a patient to enrol in the study; if both of the patient's eyes had CNV-AMD, then both were included in the analyses.	This study was conducted at 24 centers in the United States (NCT01542866).	no	16 weeks	Wet age-related macular degeneration	Distortion algorithm
Printy	2014	Cohort study, Pilot study	unclear	not reported	not reported	In a movement disorders clinic on the day of an outpatient appointment.	no	not reported	Parkinson's disease and severity	Quantification of the severity of Parkinson's motors symptoms using an application that collects kinematic data and extracts quantitative features using signal processing techniques, support vector machine classifier.
Lagido	2014	Diagnostic case-control	unclear	not reported	not reported	Samples were collected from heart failure patients at rest in Hospital S. Joao in Porto.	no	not reported	Atrial fibrillation	Detect heart rate and heart rate variability using a photoplethysmogram signal with the user's fingertip placed over the smartphone camera.

Maier	2014	Diagnostic case-control	Prospective	Images were excluded from evaluation in case of poor quality images, other elements in the image not belonging to the lesion e.g. hair, images containing more than one lesion, incomplete imaged lesions, non-melanocytic lesions, two-point differences cases (results in non-consecutive risk classes mainly due to inappropriate imaging angle or distance). The cases with an equal number of results in two consecutive risk classes, so-called tie cases (e.g. 1 high risk, 1 medium risk and 1 low risk results), were also excluded.	not reported	We included 195 melanocytic lesions in consecutive patients seen routinely for skin cancer screening at the Department of Dermatology, University Hospital of Munich, Germany after obtaining written informed consent.	yes	not reported	Melanoma	Risk assessment algorithm
Ellis	2015	Diagnostic case-control with age-matched healthy controls	unclear	not reported	A telephone questionnaire was first administered to screen out potential subjects who (1) are not within the age range of 40 to 85; (2) have any problems with their hearing; (3) are not able to walk independently without an aid; (4) have joint problems or other neurological, musculoskeletal or medical problems that can affect walking; (5) have sustained a fall within the past year that continues to affect their walking pattern; (6) have had surgery to implant a device (e.g., deep brain stimulation or pacemaker). Subjects who satisfied all six criteria were invited to participate in the study. Upon arrival at the testing location, four clinical assessments were administered.	All subjects were recruited through the Singapore General Hospital clinics. For safety reasons, the inclusion/exclusion criteria for the present study precluded patients with severe gait dysfunction; most patients in the present sample would be considered to have "moderately advanced" disease. Whether SmartMOVE would perform as well in the case of severe gait dysfunction (e.g., shuffling steps or frequent gait freezing episodes) is thus unknown.	no	not reported	Parkinson's disease	Smartphone's inertial measurement unit to record gait movements during walking. / The accuracy of smartphone-based gait analysis (utilizing the smartphone's built-in tri-axial accelerometer and gyroscope to calculate successive step times and step lengths) was validated against two heel contact-based measurement devices: heel-mounted footswitch sensors (to capture step times) and an instrumented pressure sensor mat (to capture step lengths).

Ellis	2015	Diagnostic case-control with age-matched healthy controls	unclear	not reported	A telephone questionnaire was first administered to screen out potential subjects who (1) are not within the age range of 40 to 85; (2) have any problems with their hearing; (3) are not able to walk independently without an aid; (4) have joint problems or other neurological, musculoskeletal or medical problems that can affect walking; (5) have sustained a fall within the past year that continues to affect their walking pattern; (6) have had surgery to implant a device (e.g., deep brain stimulation or pacemaker). Subjects who satisfied all six criteria were invited to participate in the study. Upon arrival at the testing location, four clinical assessments were administered.	All subjects were recruited through the Singapore General Hospital clinics. For safety reasons, the inclusion/exclusion criteria for the present study precluded patients with severe gait dysfunction; most patients in the present sample would be considered to have "moderately advanced" disease. Whether SmartMOVE would perform as well in the case of severe gait dysfunction (e.g., shuffling steps or frequent gait freezing episodes) is thus unknown.	no	not reported	Healthy subjects	Smartphone's inertial measurement unit to record gait movements during walking. / The accuracy of smartphone-based gait analysis (utilizing the smartphone's built-in tri-axial accelerometer and gyroscope to calculate successive step times and step lengths) was validated against two heel contact-based measurement devices: heel-mounted footswitch sensors (to capture step times) and an instrumented pressure sensor mat (to capture step lengths).
Nishiguchi	2014	Open-label, follow-up study	unclear	Other musculoskeletal disorders, cognitive disorders, Parkinson's disease, stroke, or unable to walk unassisted over 15m using walking aids. Patients with previous surgery in the lower extremities were also excluded.	Patients with rheumatoid arthritis defined by the 1987 or 2010 American College of Rheumatology criteria were included.	not reported	no	unclear	Rheumatoid arthritis, disease activity	The modified Health Assessment Questionnaire (mHAQ), self-assessed TJC (self-assessed tender joint count, out of 49 joints), and self-assessed SJC (self-assessed swollen joint count, out of 46 joints) were recorded on the smartphone application that we developed. The mHAQ, a self-reported measure of physical function to quantify functional disability. The mHAQ is expressed on a scale ranging from 0 to 3, where 0 = no disability and 3 = severe functional disability. Gait Analysis: The participants were instructed to walk along a 15-m walkway at their preferred speed. Trunk linear accelerations were measured by participants themselves with the smartphone as they walked on the walkway. The smartphone was kept adjacent to the L3 spinous process, which is close to where the body's center of mass is believed to be located during quiet standing using a semi-elastic belt.

Shinohara	2013	Feasibility study	unclear	Other musculoskeletal disorders, cognitive disorders, Parkinson's disease, stroke, or unable to walk over 10 m unassisted. Patients with previous surgery in the lower extremities were also excluded.	Patients with rheumatoid arthritis as defined by the American College of Rheumatology 1987 or 2010 criteria were included.	The participants were patients who attended the rheumatology outpatient clinic of Kyoto University Hospital.	no	Until the next hospital visit: mean duration, 35.5 ± 11.3 days	Rheumatoid arthritis, disease activity	Linear trunk accelerations are gathered by the participants' smartphones (kept in a waist pouch) as they walked for 10 seconds at their preferred speed. Peak frequency (PF), autocorrelation peak (AC), and coefficient of variance (CV) of the acceleration peak intervals. The PF value indicates the gait cycle, which is the time taken for 1 step. The AC value indicates the degree of gait balance, so a higher AC value indicates a greater degree of balance. The CV value indicates the degree of gait variability, i.e., the variability in the elapsed time between the first contacts of 2 consecutive footfalls. The modified Health Assessment Questionnaire (mHAQ), self-assessed TJC (self-assessed tender joint count, out of 49 joints), and self-assessed SJC (self-assessed swollen joint count, out of 46 joints) were recorded on the smartphone application that we developed. The mHAQ, a self-reported measure of physical function to quantify functional disability in RA. The mHAQ is expressed on a scale ranging from 0 to 3, where 0 = no disability and 3 = severe functional disability. General health condition and pain condition were recorded on the smartphone using a visual analogue scale (VAS). Gait Analysis: The participants were instructed to walk along a 15-m walkway at their preferred speed. Trunk linear accelerations were measured by participants themselves with the smartphone as they walked on the walkway. The smartphone was kept adjacent to the L3 spinous process, which is close to where the body's center of mass is believed to be located during quiet standing using a semi-elastic belt.
-----------	------	-------------------	---------	--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------	----------------------------------------------------------------------------------------------------------------------------	--------------------------------------------------------------------------------------------------------------	----	----------------------------------------------------------------	----------------------------------------	---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------

Yamada	2011	Cross-sectional study	unclear	We excluded participants based on the following exclusion criteria: other musculoskeletal disorders, cognitive disorders, Parkinson's disease, stroke, or unable to walk unassisted over 15 m using current walking aids.	Patients with rheumatoid arthritis defined by the American College of Rheumatology 1987 criteria were included.	This was a cross-sectional study performed between April 2011 and May 2011 in the rheumatology outpatient clinics of Kyoto University Hospital. A total of 39 RA patients (mean age, 65.9 ± 10.0 years) participated.	not reported	not reported	Rheumatoid arthritis, disease activity	The smartphone used in this study includes an acceleration sensor, a recording device, and a computer program for processing the acceleration signals. Trunk linear accelerations were measured using the smartphone while the subject walked on the walkway. The smartphone was attached to the L3 spinous process using a semi-elastic belt. Before measurements, the accelerometer of the smartphone was calibrated statically against gravity. The accelerometer of the smartphone sampled at 33 Hz. The recorded signals were analysed by the application developed in the android environment. Gait analysis: The participants were instructed to walk on a 20-m walk- way at their preferred speed. All participants wore their usual walking shoes, avoiding high heels and hard-soled shoes. The mid (10-m) walking time was measured using an electronic stopwatch.
Yamada	2011	Cross-sectional study	unclear	We excluded participants based on the following exclusion criteria: other musculoskeletal disorders, cognitive disorders, Parkinson's disease, stroke, or unable to walk unassisted over 15 m using current walking aids.	not reported	Twenty older individuals also took part in this experiment as control participants.	not reported	not reported	Healthy Control	The smartphone used in this study includes an acceleration sensor, a recording device, and a computer program for processing the acceleration signals. Trunk linear accelerations were measured using the smartphone while the subject walked on the walkway. The smartphone was attached to the L3 spinous process using a semi-elastic belt. Before measurements, the accelerometer of the smartphone was calibrated statically against gravity. The accelerometer of the smartphone sampled at 33 Hz. The recorded signals were analysed by the application developed in the android environment. Gait analysis: The participants were instructed to walk on a 20-m walk- way at their preferred speed. All participants wore their usual walking shoes, avoiding high heels and hard-soled shoes. The mid (10-m) walking time was measured using an electronic stopwatch.

Lee	2013	Case-control study, with-in	Prospective	not reported	not reported	Patients who presented for electrical cardioversion to the University of Massachusetts Medical Center (UMMC) cardiac electrophysiology laboratory were recruited by trained study personnel (McManus, Mathias)	no	not reported	Atrial fibrillation	Detect Atrial fibrillation (AF) and non-sinus rhythm (NSR) using a photoplethysmogram signal with the user's fingertip placed over the smartphone camera. AF and NSR detection is based on threshold values derived from the MIT-BIH AF and MIT-BIH NSR databases using statistical method RMSSD (Root mean square of successive differences).
Lee	2013	Case-control study, with-in	Prospective	not reported	not reported	Patients who presented for electrical cardioversion to the University of Massachusetts Medical Center (UMMC) cardiac electrophysiology laboratory were recruited by trained study personnel (McManus, Mathias)	no	not reported	Atrial fibrillation	Detect Atrial fibrillation (AF) and non-sinus rhythm (NSR) using a photoplethysmogram signal with the user's fingertip placed over the smartphone camera. AF and NSR detection is based on threshold values derived from the MIT-BIH AF and MIT-BIH NSR databases using statistical method RMSSD (Root mean square of successive differences).
Lee	2013	Case-control study, with-in	Prospective	not reported	not reported	Patients who presented for electrical cardioversion to the University of Massachusetts Medical Center (UMMC) cardiac electrophysiology laboratory were recruited by trained study personnel (McManus, Mathias)	no	not reported	Atrial fibrillation	Detect Atrial fibrillation (AF) and non-sinus rhythm (NSR) using a photoplethysmogram signal with the user's fingertip placed over the smartphone camera. AF and NSR detection is based on threshold values derived from the MIT-BIH AF and MIT-BIH NSR databases using statistical method RMSSD (Root mean square of successive differences).
Zhu	2014	unclear	unclear	None of the patients had a coexisting dementia (Mini-Mental State Examination score >24 points) or other diagnosed neurological impairments.	Patients diagnosed with idiopathic Parkinson's disease from Singapore General Hospital; All patients were under stable medication regimens for the preceding four weeks, and were tested at least 30 minutes after taking morning medications.	Patients diagnosed with idiopathic Parkinson's disease were recruited from Singapore General Hospital.	no	not reported	Parkinson's disease	To help "scale up" rhythmic auditory cueing (RAC) for wider distribution, we have developed an iOS-based Rhythmic Auditory Cueing Evaluation (iRACE) mobile application to deliver RAC and assess motor performance in PD patients. The touchscreen of the mobile device is used to assess motor timing during index finger tapping, and the device's built-in tri-axial accelerometer and gyro- scope to assess step time and step length during walking. Novel machine learning-based gait analysis algorithms have been developed for iRACE, including heel strike detection, step length quantification, and left-versus-right foot identification.

Wolf	2013	Diagnostic case-control	Convenient	<p>Images that contained any identifiable features such as facial features, tattoos, or labels with patient information were either excluded or cropped to remove the identifiable features or information. Lesions with equivocal diagnoses such as "melanoma cannot be ruled out" or "atypical melanocytic proliferation" were excluded, as were Spitz nevi, pigmented spindle cell nevus of Reed, and other uncommon or equivocal lesions. We also excluded lesions with moderate or high-grade atypia given the controversy over their management. Two of the investigators then reviewed all images for image quality and omitted those that were of poor quality or resolution.</p>	<p>Images for which there was a clear histologic diagnosis rendered by a board-certified pathologist. The remaining images were stratified into one of the following categories: invasive melanoma, melanoma in situ, lentigo, benign nevus (including compound, junctional, and low-grade dysplastic nevi), dermatofibroma, seborrheic keratosis, and hemangioma. We only used close-up images of lesions.</p>	not reported	no	not reported	Melanoma	<p>Application 1 uses an automated algorithm to detect the border of the lesion, although it also allows manual input to confirm or change the detected border. It is the only application we tested that has this feature of user input for border detection. The application then analyses the image and gives an assessment of "problematic," which we considered to be a positive test, "ok," which we considered to be a negative test, or "error" if the image could not be assessed by the application. We categorized the latter group as unevaluable.</p>
Wolf	2013	Diagnostic case-control	Convenient	<p>Images that contained any identifiable features such as facial features, tattoos, or labels with patient information were either excluded or cropped to remove the identifiable features or information. Lesions with equivocal diagnoses such as "melanoma cannot be ruled out" or "atypical melanocytic proliferation" were excluded, as were Spitz nevi, pigmented spindle cell nevus of Reed, and other uncommon or equivocal lesions. We also excluded lesions with moderate or high-grade atypia given the controversy over their management. Two of the investigators then reviewed all images for image quality and omitted those that were of poor quality or resolution.</p>	<p>Images for which there was a clear histologic diagnosis rendered by a board-certified pathologist. The remaining images were stratified into one of the following categories: invasive melanoma, melanoma in situ, lentigo, benign nevus (including compound, junctional, and low-grade dysplastic nevi), dermatofibroma, seborrheic keratosis, and hemangioma. We only used close-up images of lesions.</p>	not reported	no	not reported	Melanoma	<p>Application 2 uses an automated algorithm to evaluate an image that has been uploaded by the user. The output given is either "melanoma," which we considered to be a positive test, or "looks good" which we considered to be a negative test. If the image could not be analysed a message of "skin condition not found" was given and we considered the image unevaluable.</p>

Wolf	2013	Diagnostic case-control	Convenient	Images that contained any identifiable features such as facial features, tattoos, or labels with patient information were either excluded or cropped to remove the identifiable features or information. Lesions with equivocal diagnoses such as "melanoma cannot be ruled out" or "atypical melanocytic proliferation" were excluded, as were Spitz nevi, pigmented spindle cell nevus of Reed, and other uncommon or equivocal lesions. We also excluded lesions with moderate or high-grade atypia given the controversy over their management. Two of the investigators then reviewed all images for image quality and omitted those that were of poor quality or resolution.	Images for which there was a clear histologic diagnosis rendered by a board-certified pathologist. The remaining images were stratified into one of the following categories: invasive melanoma, melanoma in situ, lentigo, benign nevus (including compound, junctional, and low-grade dysplastic nevi), dermatofibroma, seborrheic keratosis, and hemangioma. We only used close-up images of lesions.	not reported	no	not reported	Melanoma	Application 3 asks the user to upload an image to the application and then to position it within a box to ensure that the correct lesion is analysed. The output given by the application is "high risk," which we considered to be a positive test, or "medium risk" or "low risk," both of which we considered to be a negative test. The presence of a medium risk category in this application presented some difficulty in analysis as it was the only application tested that gave an intermediate output. Thus, we did perform sensitivity and specificity analysis with "medium risk" lesions counting as a positive test as well since it is not clear how a user would interpret such a result. Some lesions generated a message of "error" and these were considered unevaluable.
McManus	2013	Diagnostic case-control, within	Prospective	not reported	not reported	In a prospectively recruited cohort of participants undergoing cardioversion for atrial fibrillation	no	not reported	Atrial fibrillation	Mean square of successive differences (RMSSD): Our application acquired pulsatile signals by illuminating the fingertip using the standard iPhone lamp and recording video signal (30 frames/s) for 2 minutes. The signal was processed by averaging 50x50 green band pixels per frame. We interpolated the pulsatile signal to 30 Hz using a cubic spline algorithm followed by peak detection. As described in prior work, we use a peak detection algorithm that uses a filter bank with estimates of heart rate, variable cut-off frequencies, rank-order nonlinear filters, and decision logic as well as motion noise correction.

McManus	2013	Diagnostic case-control, within	Prospective	not reported	not reported	In a prospectively recruited cohort of participants undergoing cardioversion for atrial fibrillation	no	not reported	Atrial fibrillation	Shannon entropy (ShE): Our application acquired pulsatile signals by illuminating the fingertip using the standard iPhone lamp and recording video signal (30 frames/s) for 2 minutes. The signal was processed by averaging 50x50 green band pixels per frame. We interpolated the pulsatile signal to 30 Hz using a cubic spline algorithm followed by peak detection. As described in prior work, we use a peak detection algorithm that uses a filter bank with estimates of heart rate, variable cut-off frequencies, rank-order nonlinear filters, and decision logic as well as motion noise correction.
McManus	2013	Diagnostic case-control, within	Prospective	not reported	not reported	In a prospectively recruited cohort of participants undergoing cardioversion for atrial fibrillation	no	not reported	Atrial fibrillation	Combined statistical method: root mean square of successive RR difference (RMSSD/mean) and Shannon entropy (ShE): Our application acquired pulsatile signals by illuminating the fingertip using the standard iPhone lamp and recording video signal (30 frames/s) for 2 minutes. The signal was processed by averaging 50x50 green band pixels per frame. We interpolated the pulsatile signal to 30 Hz using a cubic spline algorithm followed by peak detection. As described in prior work, we use a peak detection algorithm that uses a filter bank with estimates of heart rate, variable cut-off frequencies, rank-order nonlinear filters, and decision logic as well as motion noise correction.

Takuya	2015	Cross-sectional study	Convenient	Exclusion criteria were (1) severe cardiovascular, respiratory, musculoskeletal, or neurologic disorder other than stroke that affected gait performance; (2) unable to understand the instructions because of communication problem or moderate to severe cognitive dysfunction (i.e., 5 or more errors on the Short Portable Mental Status Questionnaire [SPMSQ]); and (3) household ambulators walked only indoors or only mobilized during rehabilitation sessions.	Inclusion criteria were (1) more than 12 months since stroke onset and (2) ability to walk 16 m independently with or without a single point cane and/or an orthosis.	Community-dwelling adults with chronic stroke receiving day care services were recruited and screened for inclusion and exclusion criteria. conducted in 2 day care centers for elderly adults in Saitama, Japan	no	not reported	Chronic stroke, falls	Gait characteristics
Takuya	2015	Cross-sectional study	Convenient	Exclusion criteria were (1) severe cardiovascular, respiratory, musculoskeletal, or neurologic disorder other than stroke that affected gait performance; (2) unable to understand the instructions because of communication problem or moderate to severe cognitive dysfunction (ie, 5 or more errors on the Short Portable Mental Status Questionnaire [SPMSQ]); and (3) household ambulators walked only indoors or only mobilized during rehabilitation sessions.	Inclusion criteria were (1) more than 12 months since stroke onset and (2) ability to walk 16 m independently with or without a single point cane and/or an orthosis.	Community-dwelling adults with chronic stroke receiving day care services were recruited and screened for inclusion and exclusion criteria. conducted in 2 day care centers for elderly adults in Saitama, Japan	no	not reported	chronic stroke, no falls	Gait characteristics
Chadwick	2014	Case-control study	Convenient	not reported	not reported	The first step in the app testing was the selection of 15 high-quality, full resolution, digital images (.jpg) of melanocytic skin lesions of varying risk (5 melanomas, 10 benign nevi) from the image archive of the study dermatology expert (HPS) to assess the accuracy of the app analysis software.	no	not reported	Melanoma	SkinScan - Asymmetry/Border: Analysis by inbuilt pattern recognition software. / Color: Analysis by input comparison algorithms. / Diameter: no input. / Evolution: no input

Chadwick	2014	Case-control study	Convenient	not reported	not reported	The first step in the app testing was the selection of 15 high-quality, full resolution, digital images (.jpg) of melanocytic skin lesions of varying risk (5 melanomas, 10 benign nevi) from the image archive of the study dermatology expert (HPS) to assess the accuracy of the app analysis software.	no	not reported	Melanoma	MelApp - Asymmetry/Border: Analysis by inbuilt pattern recognition software. Area could be limited to focus the analysis. / Color: Analysis by input comparison algorithms. / Diameter: manual sliding scale input. / Evolution: manual sliding scale input
Chadwick	2014	Case-control study	Convenient	not reported	not reported	The first step in the app testing was the selection of 15 high-quality, full resolution, digital images (.jpg) of melanocytic skin lesions of varying risk (5 melanomas, 10 benign nevi) from the image archive of the study dermatology expert (HPS) to assess the accuracy of the app analysis software.	no	not reported	Melanoma	Mole Detective - Asymmetry/Border: Analysis by inbuilt pattern recognition software. / Color: Analysis by input comparison algorithms. / Diameter: Manual input of <6mm, -mm, or >6mm. / Evolution: No input for analysis. Reminder can be set for future use.
Chadwick	2014	Case-control study	Convenient	not reported	not reported	The first step in the app testing was the selection of 15 high-quality, full resolution, digital images (.jpg) of melanocytic skin lesions of varying risk (5 melanomas, 10 benign nevi) from the image archive of the study dermatology expert (HPS) to assess the accuracy of the app analysis software.	no	not reported	Melanoma	Spot Mole Plus - Asymmetry/Border: Analysis by inbuilt pattern recognition software. Manual adjustment of lesion border available/ Color: Analysis by input comparison algorithms. / Diameter: Manual input of numeric value. / Evolution: no input for past history of change. Can perform serial analysis of lesion images.
Chadwick	2014	Case-control study	Convenient	not reported	not reported	The first step in the app testing was the selection of 15 high-quality, full resolution, digital images (.jpg) of melanocytic skin lesions of varying risk (5 melanomas, 10 benign nevi) from the image archive of the study dermatology expert (HPS) to assess the accuracy of the app analysis software.	no	not reported	Melanoma	Dr. Mole Premium - Asymmetry/Border: Analysis by inbuilt pattern recognition software. Comparison of lesion quadrants for asymmetry./ Color: Analysis by input comparison algorithms. / Diameter: Manual sliding scale input. / Evolution: manual input of "none", "slow" & "fast" with time frames for each.

Winther	2015	Longitudinal Pilot study	Convenient	Exclusion criteria were the bilateral presence of currently inactive lesions judged to run a small risk of recurrence, the presence of additional, non- maculopathic causes of visual loss, and inability to participate in conventional acuity testing. Twenty-eight patients partook in the study. Those who had the same type of lesions in both eyes (active or inactive) provided results from the least involved eye only whereas those who had different types of lesions provided results from both eyes.	not reported	Patients were recruited from the wet age-related macular degeneration treatment programme of the Retina Unit at the Sahlgrenska University Hospital, a tertiary-care center.	no	Average monitoring of average of 30 weeks. Another time they report an average of 39.	Wet age-related macular degeneration	For formal analysis, all MBT plots were carefully evaluated by subjective inspection, epoch by epoch. Epochs showing trends of decreasing scores, or increasing variation, or both, were rated worse. Epochs showing the opposite evolution were rated better. All other epochs were rated stable. ETDRS results were rated similarly, using direct numerical comparisons of scores. The outcomes of the clinical examinations, which included biomicroscopy and scrutiny of OCT parameters and maps, were summarized in the same manner.
Winther	2015	Longitudinal Pilot study	Convenient	Exclusion criteria were the bilateral presence of currently inactive lesions judged to run a small risk of recurrence, the presence of additional, non- maculopathic causes of visual loss, and inability to participate in conventional acuity testing. Twenty-eight patients partook in the study. Those who had the same type of lesions in both eyes (active or inactive) provided results from the least involved eye only whereas those who had different types of lesions provided results from both eyes.	not reported	Twenty control subjects were recruited primarily from patients' relatives or other accompanying persons.	no	Average monitoring of average of 30 weeks. Another time they report an average of 39.	Healthy relatives or accompanying	not reported

Wang	2013	Cross-sectional study	Convenient	not reported	1) AMD or DR with corrected Early Treatment of Diabetic Retinopathy Study (ETDRS) VA of 20/100 or better in at least one eye, 2) ophthalmic evaluation by retina specialists with clinical and spectral-domain (SD)-OCT documentation, 3) no retinal pathology other than AMD or DR, 4) no concurrent systemic illness affecting the retina, and 5) no dementia or other limitation that would prevent the patient from performing a self-test of visual function. Patients with AMD and DR were recruited at various disease stages, including those under active anti-vascular endothelial growth factor treatment. Patients with epiretinal membrane or pigment epithelial detachment were not excluded.	Patients with AMD and DR were recruited from the clinic of the Department of Ophthalmology, UT Southwestern Medical Center.	no	not reported	Age-related macular degeneration
------	------	-----------------------	------------	--------------	-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------	-----------------------------------------------------------------------------------------------------------------------------	----	--------------	----------------------------------

Wang	2013	Cross-sectional study	unclear	not reported	<p>1) AMD or DR with corrected Early Treatment of Diabetic Retinopathy Study (ETDRS) VA of 20/100 or better in at least one eye, 2) ophthalmic evaluation by retina specialists with clinical and spectral-domain (SD)-OCT documentation, 3) no retinal pathology other than AMD or DR, 4) no concurrent systemic illness affecting the retina, and 5) no dementia or other limitation that would prevent the patient from performing a self-test of visual function. Patients with AMD and DR were recruited at various disease stages, including those under active anti-vascular endothelial growth factor treatment. Patients with epiretinal membrane or pigment epithelial detachment were not excluded.</p>	<p>Patients with AMD and DR were recruited from the clinic of the Department of Ophthalmology, UT Southwestern Medical Center.</p>	no	not reported	Diabetic retinopathy
------	------	-----------------------	---------	--------------	--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------	------------------------------------------------------------------------------------------------------------------------------------	----	--------------	----------------------

Wang	2013	Cross-sectional study	unclear	not reported	1) AMD or DR with corrected Early Treatment of Diabetic Retinopathy Study (ETDRS) VA of 20/100 or better in at least one eye, 2) ophthalmic evaluation by retina specialists with clinical and spectral-domain (SD)-OCT documentation, 3) no retinal pathology other than AMD or DR, 4) no concurrent systemic illness affecting the retina, and 5) no dementia or other limitation that would prevent the patient from performing a self-test of visual function. Patients with AMD and DR were recruited at various disease stages, including those under active anti-vascular endothelial growth factor treatment. Patients with epiretinal membrane or pigment epithelial detachment were not excluded.	Healthy subjects were recruited from the normal subject database of the Retina Foundation of the Southwest.	no	not reported	Healthy senior volunteers	
Woods	2014	unclear	unclear	not reported	not reported	Participants diagnosed with Parkinson's disease	no	unclear	Parkinson tremor	Smartphone application that uses discrete wavelet transforms and support vector machines to discriminate between Parkinson's and Essential postural tremors / Triaxial, digital acceleration sensor. The 6 experimental tasks were: (1) tremor with eyes open (Vis+); (2) tremor with eyes closed (Vis-); (3) tremor while attending to the active tremor hand (Bubble); (4) tremor while attending to a laser target at 2 m (Laser2); (5) tremor while attending to a laser target at 1 m (Laser1); (6) tremor while not attending to the hand but while counting backwards by 3 (Counting).

Woods	2014	unclear	unclear	not reported	not reported	Participants with essential tremor	no	unclear	Essential Tremor	Smartphone application that uses discrete wavelet transforms and support vector machines to discriminate between Parkinson's and Essential postural tremors / Triaxial, digital acceleration sensor. The 6 experimental tasks were: (1) tremor with eyes open (Vis+); (2) tremor with eyes closed (Vis-); (3) tremor while attending to the active tremor hand (Bubble); (4) tremor while attending to a laser target at 2 m (Laser2); (5) tremor while attending to a laser target at 1 m (Laser1); (6) tremor while not attending to the hand but while counting backwards by 3 (Counting).
Kostikis	2014	Pilot study	Convenient	not reported	not reported	Subjects participating in this study were all Parkinson's disease patients recruited from the outpatient clinic of the 1st Department of Neurology at the Aristotle University of Thessaloniki.	no	not reported	Parkinson's Tremor	Using a smartphone-based platform, which processes the phone's accelerometer and gyroscope signals to detect and measure hand tremor. / In this work we are initially interested in resting tremor so we asked the subjects to "wear" an iPhone (fitted on a glove as in [3]) on top of their hand while sitting in a chair comfortably and resting both their hands on their lap, keeping that position for 30 seconds. The device was mounted on both their hands alternately, and each test was repeated twice for each subject.

Kostikis	2015	Cross-sectional study	Convenient	not reported	not reported	We recruited patients from the outpatient clinic of the first Department of Neurology at the Aristotle University of Thessaloniki. They all agreed to participate after they were offered a detailed explanation of the study's procedure and goals. All of them were right-handed, under L-DOPA treatment and suffering from Parkinson for more than two years.	no	not reported	Parkinson's Tremor	We attached an iPhone on our volunteer's hands using the same custom-made mounting glove]. It consists of a perforated case into which the phone "locks," and a wrist-supporting glove, both commercially available. The glove fits tightly on the volunteer's hand and the case is tightly sewn on the glove using nonelastic thread, ensuring the stability of the device on top of the hand. With the device attached, each participant had to maintain each of two prescribed postures for 30 s, while acceleration and gyroscope data were recorded by the phone. The two postures we used were the same ones used during the clinical evaluation, 1) "Extended," i.e., seated with both hands extended in front of the torso (Postural Tremor of the Hands, component 3.15 of the MDS-UPDRS) and 2) "Rest," i.e., seated with both hands placed on the arms of the chair (Rest Hand Tremor, component 3.17 of the MDS-UPDRS). The procedure was then repeated for the subject's other hand, in the same two postures. In the following, we will specify the combination of a patient's hand (Right of Left upper extremity) during each position as rR, rL, eR, and eL for rest-right, rest-left, extended-right, and extended-left, respectively.
----------	------	-----------------------	------------	--------------	--------------	------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------	----	--------------	--------------------	----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------

Kostikis	2015	Cross-sectional study	Random	They were screened for several health conditions which could exclude them from the study, such as hypertension or any movement disorder. They were also notified of the procedure and the purpose of the study before agreeing to participate.	not reported	The control group for the study, contains healthy volunteers, none of whom suffered from a movement disorder, hypertension, or diabetes.	no	not reported	Age-matched healthy volunteers	We attached an iPhone on our volunteer's hands using the same custom-made mounting glove]. It consists of a perforated case into which the phone "locks," and a wrist-supporting glove, both commercially available. The glove fits tightly on the volunteer's hand and the case is tightly sewn on the glove using nonelastic thread, ensuring the stability of the device on top of the hand. With the device attached, each participant had to maintain each of two prescribed postures for 30 s, while acceleration and gyroscope data were recorded by the phone. The two postures we used were the same ones used during the clinical evaluation, 1) "Extended," i.e., seated with both hands extended in front of the torso (Postural Tremor of the Hands, component 3.15 of the MDS-UPDRS) and 2) "Rest," i.e., seated with both hands placed on the arms of the chair (Rest Hand Tremor, component 3.17 of the MDS-UPDRS). The procedure was then repeated for the subject's other hand, in the same two postures. In the following, we will specify the combination of a patient's hand (Right of Left upper extremity) during each position as rR, rL, eR, and eL for rest-right, rest-left, extended-right, and extended-left, respectively.
----------	------	-----------------------	--------	------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------	--------------	------------------------------------------------------------------------------------------------------------------------------------------	----	--------------	--------------------------------	----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------

Kostikis	2015	Cross-sectional study	Convenient	not reported	not reported	We recruited patients from the outpatient clinic of the first Department of Neurology at the Aristotle University of Thessaloniki. They all agreed to participate after they were offered a detailed explanation of the study's procedure and goals. All of them were right-handed, under L-DOPA treatment and suffering from Parkinson for more than two years.	no	not reported	De novo Parkinson's disease: During the study, two of them were hospitalized overnight so that they could be tested in the morning before they received their medication, to approximate de novo patients.	We attached an iPhone on our volunteer's hands using the same custom-made mounting glove]. It consists of a perforated case into which the phone "locks," and a wrist-supporting glove, both commercially available. The glove fits tightly on the volunteer's hand and the case is tightly sewn on the glove using nonelastic thread, ensuring the stability of the device on top of the hand. With the device attached, each participant had to maintain each of two prescribed postures for 30 s, while acceleration and gyroscope data were recorded by the phone. The two postures we used were the same ones used during the clinical evaluation, 1) "Extended," i.e., seated with both hands extended in front of the torso (Postural Tremor of the Hands, component 3.15 of the MDS-UPDRS) and 2) "Rest," i.e., seated with both hands placed on the arms of the chair (Rest Hand Tremor, component 3.17 of the MDS-UPDRS). The procedure was then repeated for the subject's other hand, in the same two postures. In the following, we will specify the combination of a patient's hand (Right of Left upper extremity) during each position as rR, rL, eR, and eL for rest-right, rest-left, extended-right, and extended-left, respectively.
----------	------	-----------------------	------------	--------------	--------------	------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------	----	--------------	------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------	----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------

Arora	2015	Pilot study	Convenient	Importantly, this study did not include individuals with other parkinsonian or tremor disorders that may be more difficult to differentiate from Parkinson's disease.	not reported <sup>9</sup>	Individuals with Parkinson diagnosed clinically by a movement disorder specialist, were recruited from an academic movement disorder clinic (Johns Hopkins) and all participants provided informed consent.	no	1month / study duration: average 34.4 days	Parkinson's disease	(1) (voice test) say the sustained phonation 'aaah' for as long and as steadily as possible; (2) (posture test) stand upright unaided for thirty seconds; (3) (gait test) walk twenty steps forward, turn around, and return back to the starting position; (4) (finger tapping test) tap the screen alternately keeping a regular rhythm; and (5) (reaction time test) press and hold the on-screen button as soon as it appears and release it as soon as it disappears. The participants were asked to conduct the above specified tests four times daily: just before taking their first (morning) dose of levodopa (or in one case, rasagiline), one hour later, mid- afternoon, and before going to bed.
Arora	2015	Pilot study	Convenient	Known neurological disorder	not reported	Were recruited from an academic movement disorder clinic (Johns Hopkins) and all participants provided informed consent. Control participants were spouses, caregivers, relatives, or colleagues of an individual with PD	no	1month / study duration: average 34.4 days	Healthy volunteers (Control participants)	(1) (voice test) say the sustained phonation 'aaah' for as long and as steadily as possible; (2) (posture test) stand upright unaided for thirty seconds; (3) (gait test) walk twenty steps forward, turn around, and return back to the starting position; (4) (finger tapping test) tap the screen alternately keeping a regular rhythm; and (5) (reaction time test) press and hold the on-screen button as soon as it appears and release it as soon as it disappears. The participants were asked to conduct the above specified tests four times daily: just before taking their first (morning) dose of levodopa (or in one case, rasagiline), one hour later, mid- afternoon, and before going to bed.
Wadhawan	2011	Case-control	Convenient	Image artefacts	not reported	In this study, we use images from a large library of skin cancer images that were uploaded to the phone.	not reported	not reported	Melanoma	A support vector machine (SVM) is trained using a subset (training set) of the total images available, and the resulting classifier is used to determine whether the rest of the images (test set) are malignant or benign.

Wadhawan	2011	Case-control	Convenient	Image artefacts	not reported	In this study, we used images from a large commercial library of skin cancer images annotated by expert dermatologists that were uploaded to the phone. Intra-observer and inter-observer agreement could be low for certain criteria. To demonstrate the feasibility of our automated system, we only chose images considered as low difficulty by the experts. There were 385 low difficulty images in the database and our segmentation methods could provide a satisfactory boundary for 347 (90.13%) of them.	not reported	not reported	Melanoma	A skin lesion image can be acquired using the smartphone camera (with or without an external attachment which can provide illumination and magnification) or can be loaded from the photo library to provide the diagnosis in real time. To identify a region of interest (ROI), an image is first converted to greyscale, and then fast median filtering for noise removal is performed, and followed by ISODATA segmentation, and several morphological operations.
Daneault	2012	Observational	Random	not reported	not reported	not reported	not reported	not reported	Tremor in Essential tremor, Parkinson's disease and multiples Sclerosis	Algorithms for tremor recording and online analysis
Doukas	2012	Case-Control	Convenient	not reported	not reported	For the initial evaluation of the system a dataset consisting of over 3000 skin lesion image sets of manually classified images has been utilized. The dataset contained about 800 images with melanoma, 600 with dysplastic nevus and the rest 1600 images with benign nevi.	not reported	not reported	Melanoma	SVM algorithm / automated skin lesion assessment system based on mobile technologies that can be used by patients for an early characterization of lesions and estimation for further assessment.
Doukas	2012	Usability Questionnaire	Convenient	not reported	not reported	For the initial evaluation of the system a dataset consisting of over 3000 skin lesion image sets of manually classified images has been utilized. The dataset contained about 800 images with melanoma, 600 with dysplastic nevus and the rest 1600 images with benign nevi.	not reported	not reported	Melanoma	SVM algorithm / automated skin lesion assessment system based on mobile technologies that can be used by patients for an early characterization of lesions and estimation for further assessment.
Fuadah	2015	Cross-sectional study	Convenient	not reported	not reported	In this research, we use 160 eyes images. We divided them into two categories: cataract (80 images) and normal (80 images). The training data set consist of 40 normal image and 40 cataract images.	not reported	not reported	Cataract	The interface of MCataract application consists of main activity and start activity that has functions, taking a picture or loading image from the gallery, cropping to obtain the pupil area as RoI, showing the pupil area obtained, and diagnosing the condition of eye image. K-Nearest Neighbor (k-NN) as classification method.

Do	2014	Cross-sectional study	Convenient	not reported	not reported	The database includes 81 color images provided by National Skin Center, Singapore	not reported	not reported	Melanoma	In this paper, we proposed (i) an efficient segmentation scheme by combining fast skin detection and fusion of two fast segmentation results; (ii) new features which efficiently capture the color variation and border irregularity of segmented lesion and (iii) an efficient criterion for selecting features. From selected features by proposed criterion, an automatic melanoma diagnosis system using mobile is developed.
Ramlakhan	2011	Case-control study	Convenient	not reported	not reported	As the lesion image dataset, 37 images of benign skin lesions, and 46 images of malignant lesions were obtained from various Internet sites.	not reported	not reported	Melanoma	The system consists of three major components: image segmentation, feature calculation, and classification. It is designed to run on a mobile device with a camera, such as a smartphone or a tablet PC. A skin lesion image is converted to a monochrome image for outline contour detection. Color and shape features of the lesion are extracted and used as input to a kNN classifier.
Raknim	2016	Case study	Convenient	not reported	not reported	not reported	not reported	1 year of data collection.	Parkinson's disease / changes in the walking patterns of PD patients	PDR-based method to continuously monitor and record the patient's gait characteristics using a smartphone / identify changes in the walking patterns of a patient
Arora	2011	Cohort study	Convenient	not reported	not reported	Individuals with Parkinson's diagnosed clinically by a movement disorder specialist and control participants were recruited from an academic movement disorder clinic (Johns Hopkins).	not reported	1 month	Parkinson's disease	Using tri-axial accelerometry data for self-administered tests of gait and postural sway recorded via consumer-grade smartphones to accurately distinguish Parkinson's disease participants from healthy controls.
Arora	2011	Cohort study	Convenient	not reported	not reported	Individuals with Parkinson's diagnosed clinically by a movement disorder specialist and control participants were recruited from an academic movement disorder clinic (Johns Hopkins).	not reported	1 month	Age- and gender-matched participants	Using tri-axial accelerometry data for self-administered tests of gait and postural sway recorded via consumer-grade smartphones to accurately distinguish Parkinson's disease participants from healthy controls.

Garca	2014	unclear	unclear	not reported	not reported	All the PD patients performed the tests in Hospital S. João, Oporto, Portugal, while they were waiting for the medical consult. Only the ones indicated by a neurologist helped in this project since a few requisites were necessary (motor capabilities and no medication related symptoms).	not reported	not reported	Parkinson's disease	The smartphone application has four different components: two of them require active interaction of the patient with the smartphone (spiral and tap analysis), one requires passive interaction (gait analysis) and the last component is used by the health care professional (simple questions with simple observation skills).
Garca	2014	unclear	unclear	not reported	not reported	The control group was assembled by visiting social centers or inviting seniors to Fraunhofer's facilities.	not reported	not reported	Healthy controls	The smartphone application has four different components: two of them require active interaction of the patient with the smartphone (spiral and tap analysis), one requires passive interaction (gait analysis) and the last component is used by the health care professional (simple questions with simple observation skills).