Evaluation Studies of Robotic Rollators by the User Perspective: A Systematic Review

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1 Abstract

Background: Robotic rollators enhance the basic functions of established devices by technically advanced physical, cognitive, or sensory support to increase autonomy in persons with severe impairment. In the evaluation of such Ambient Assisted Living solutions, both the technical and user perspectives are important to prove usability, effectiveness, and safety, and to ensure adequate device application.

Objective: The aim of this systematic review is to summarize the methodology of studies evaluating
robotic rollators with focus on the user perspective and to give recommendations for future evaluation
studies.

10 Methods: A systematic literature search up to December 31, 2014 was conducted based on the 11 Cochrane Review methodology using the electronic databases PubMed and IEEE Xplore. Articles 12 were selected according to the following inclusion criteria: Evaluation studies of robotic rollators 13 documenting human-robot interaction, no case reports, published in English language.

Results: Twenty-eight studies were identified that met the predefined inclusion criteria. Large heterogeneity in the definitions of the target user group, study populations, study designs, and assessment methods was found across the included studies. No generic methodology to evaluate robotic rollators could be identified. We found major methodological shortcomings related to insufficient sample descriptions and sample sizes, and lack of appropriate, standardized and validated assessment methods. Long-term use in habitual environment was also not evaluated.

Conclusions: Apart from the heterogeneity, methodological deficits in most of the identified studies became apparent. Recommendations for future evaluation studies include: clear definition of target user group, adequate selection of subjects, inclusion of other assistive mobility devices for comparison, evaluation of the habitual use of advanced prototypes, adequate assessment strategy with established, standardized and validated methods, and statistical analysis of study results. Assessment strategies may additionally focus on specific functionalities of the robotic rollators allowing an individually tailored assessment of innovative features to document their added value.

27

28 Key words

29 Systematic review, Evaluation studies, Ambient assisted living, Robotics, Rollator, Walker, Self-help

30 devices, Human-robot interaction, Mobility, User experience

31 Introduction

32 In older persons, the ability to move independently represents a hallmark of autonomous living [1] 33 and quality of life [2], while being physically active is associated with numerous positive health 34 outcomes [3, 4]. However, sensory, motor or cognitive impairments restrict mobility in frail, older 35 persons [5]. Motor key functions such as standing, walking, or transfers are substantial challenges for 36 their daily activities leading to high risk exposure of falls as documented in residents of senior homes [6]. Effects of motor impairment are augmented by sensory deficits such as visual impairment, leading 37 38 to restricted functional independence [7], or by cognitive impairment, leading to spatio-temporal 39 disorientation or executive dysfunction [8]. To overcome or compensate for such impairments and to 40 improve the quality of life of affected persons, assistive devices as in walking aids (e.g. canes, 41 walkers, rollators) have been developed with an early focus on mobility support. They provide support 42 of postural stability and mobility [9], reduce risk of falling [10], and improve activity and participation 43 [11]. However, such conventional mobility devices may not cover the needs of persons suffering from 44 major functional or cognitive impairments.

45 In the context of Ambient Assisted Living (AAL), robotically augmented rollators with various 46 high-tech functionalities have been developed to provide physical, sensory and cognitive assistance, 47 and/or health monitoring for further support [12]. The development and evaluation of such a robotic 48 rollator (RR) is still a new, emerging research field mainly driven by technical engineering goals. 49 However, as technical functionalities translate into assistive devices for use of the target population, 50 for which these have been developed, the human-robot interaction and user perspective shifts in the 51 development focus. Apart from the sheer technical evaluation of concepts and functionalities, needs, 52 requirements, and preferences of potential users will have to guide the development and evaluation of 53 assistive technology devices [13, 14]. In addition to technical testing, which verifies the functional 54 capability of devices, an evaluation with focus on user performance, physical demands, and subjective 55 experiences of the RR is essential to prove the usability, ensure safety, and demonstrate the added 56 value for the intended user group. The change from technical to user perspective may, however, lead 57 to specific methodological challenges including the study design and assessment strategy. To our 58 knowledge, no systematic review on the evaluation of RRs with focus on the user perspective has been 59 published. Therefore, the aim of this systematic review was to summarize the methodology of studies 60 evaluating the human-robot interaction from a user perspective and to give recommendations for 61 future evaluation studies.

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63 Methods

Initial search terms were compiled and iteratively refined by team members with expertise in the clinical and in the technical research field. The literature search was conducted using the electronic databases PubMed and IEEE Xplore. Search terms included both controlled vocabulary (i.e. MeSH Terms, IEEE Terms) and keywords of relevance identified during searches. The detailed search strategy used in PubMed, which was modified for IEEE Xplore, is presented in the onlinesupplementary table 1.

70 Manual searches were performed to identify additional studies by scanning reference lists of 71 relevant articles and by reviewing key authors' own databases. Studies were searched with focus on 72 the evaluation of a RR (or robotic wheeled walker) by experiments, trials, or interventions in human 73 beings independent of the type of outcome measurement. No restrictions regarding age or health status 74 of the subjects were made. Single case reports were excluded. For the purpose of this review the term 75 'robotic' includes the normal function of a rollator enhanced by additional physical, sensory, or cognitive robotic support while walking, also including sit-to-stand transfers. Studies evaluating solely 76 77 monitoring functionalities without taking into account any user supporting functionalities or the 78 subjective user experience were excluded. The search was limited to articles in the English language 79 published up to December 31, 2014.

80 The selection process was conducted following the methodology as described in the method 81 guidelines of the Cochrane Collaboration [15]. Titles and abstracts were identified by the standardized 82 search strategy. For abstracts which met the inclusion criteria or for those with unclear status, full-text 83 articles were analyzed for inclusion. Each step of study selection, based on predefined eligibility 84 criteria, was performed independently by two reviewers (PU, CW). Any disagreements were resolved 85 by consensus or third-party adjudication (KH). After inclusion, data on the user group, sample characteristics, and the methodological approach were extracted by one researcher (CW) and 86 87 confirmed by two other researchers (PU, DS). If an article described more than one study, the results 88 for each study were extracted separately.

89

90 **Results**

91 A total of 8989 articles were identified through database searching, and another 79 were added 92 through manual searches. After removing duplicates, the initial search resulted in 8876 articles. Of 93 these, 235 were found to be related to the search topic based on title and abstract. After reviewing full 94 texts, 148 articles were excluded as they did not meet the predefined inclusion criteria (Fig. 1). 95 Another 63 were discarded, as these articles described either identical experiments with the same RR, 96 or various stages of development of a certain RR. In both cases, the article providing the most 97 comprehensive information with focus on the user perspective was included. If different articles 98 contained similar information, the one with the most recent development stage was included. Twenty-99 four articles published between 2001 and 2015 were identified for inclusion in the review. As two 100 articles reported on two [16, 17] and one article on three independent studies [18], the final data 101 extraction was based on 28 studies[†]. The detailed review results extracted for each study are presented

[†] When necessary, the individual studies of these articles are distinguished with numeric coding (i.e. [16^{1,2}], [17^{1,2}], [18^{1,2,3}])

in the online supplementary table 2, containing information on the names of devices, the definition ofuser groups, study sample, study object, study design, and selected assessment methods.

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(*Please insert figure 1 about here*) **Fig. 1:** Flow chart of the study selection process and extraction methodology

108 User Group Definitions

109 Apart from two articles [19, 20], all mentioned a target user group for the RR; however, their 110 definition differed substantially in accuracy and explicitness. Five articles provided a generic 111 description in broad terms such as 'elderly (disabled) people' [21-25], two defined users by setting-112 specific characteristics such as 'persons in nursing and assisted living homes', partly amended by 113 disease-related criteria (e.g. Alzheimer's disease, stroke) [26, 27], and ten provided brief information 114 on users' motor-functional (e.g. 'with mobility problems'), cognitive (e.g. 'with cognitive 115 impairment') and/or visual status (e.g. 'visually impaired') [17, 18, 28-35], but without staging 116 impairment levels based on any screening or assessment instrument. Three articles described users by 117 disease categories (e.g. Parkinson's disease, hemiplegia) [16, 36, 37] without detailed information on 118 the patients' functional impairment level. Specific impairment-related definitions based on established, 119 validated assessment methods (i.e. Walking Index for Spinal Cord Injury [WISCI II], Functional 120 Ambulation Classification) were documented in only two articles [12, 38].

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122 Study Samples

123 The mean sample size of studies was 7.2 (standard deviation [SD] \pm 4.3). The exact number of 124 subjects was not reported in five studies [18^{1,2,3}, 35, 37]. No study presented a sample size calculation. 125 Samples differed considerably regarding age, impairments, or diseases. The age of subjects ranged

126 from 14 [22] to 97 years [31] with age information lacking in half of the studies (14 of 28) [16¹, 17¹, 127 $18^{1,2,3}$, 20, 23, 25, 27-29, 35, 37].

Thirteen studies included subjects with motor, functional, cognitive, visual and/or neurological 128 impairments [12, 16^{1,2}, 17^{1,2}, 26, 27, 30-32, 34, 36, 38], whereas a convenient (e.g. 'ordinary adult 129 males') [19, 20, 23, 24, 33], mixed (e.g. 'healthy subjects and subjects with motor and cognitive 130 impairment') [18^{1,2,3}, 21, 22, 29, 35, 37] or setting-specific sample (e.g. 'residents of retirement 131 132 facility') [28] was used in 14 studies. In studies including impaired subjects, definitions and staging of the severity level of impairment were mostly absent (15 of 20) [17^{1,2}, 18^{2,3}, 22, 26, 29-32, 34, 35, 37, 133 134 38]. In only six studies, motor-functional or cognitive impairment levels were defined by established 135 and validated screening or assessment instruments (e.g. Timed up and Go [TUG], Mini-Mental State Examination) [12, 16^{1,2}, 21, 27, 36]. 136

In ten studies, subjects did not match with the predefined user group [18^{1,2,3}, 22-24, 27, 28, 33, 37].
However, due to the unspecific and wide-ranging user group definitions given in a number of articles,
most studies (15 of 28) were carried out with subjects who were covered by these broad definitions

[12, 16^{1,2}, 17^{1,2}, 21, 26, 29-32, 34-36, 38]. In three studies, a user group definition and/or a description
of the study sample was completely missing [19, 20, 25].

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143 **Design of Studies**

Depending on study objectives, three different types of studies were performed: (1) observational
studies; (2) comparative studies, or (3) interventional studies.

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147 Observational Studies

Fourteen articles reported on observational studies [12, 18, 20, 22, 24, 29, 35, 37] or single observational experiments as part of their studies [16, 17, 23, 26, 28, 33], focusing predominantly on the verification of technical capability and/or the subjective user evaluation of RR. User performance was used as the study object in only one of these studies [26]. In observational studies/experiments, outcomes were only descriptively presented, without providing any reference values.

153

154 *Comparative Studies*

155 Fourteen articles included comparative studies [19, 21, 25, 27, 28, 30-32, 34, 38] or single comparative experiments in addition to observations [16, 17, 26, 33]. Comparisons were further 156 157 distinguished into four categories: (1) 'inter-device comparisons' in which RR and conventional 158 devices (e.g. cane, folding/wheeled walker) or fully unassisted walking/sit-to-stand transfers were 159 compared [19, 21, 26, 27, 30, 32, 34, 38]; (2) 'intra-device comparisons' in which different assistance 160 levels (e.g. activated vs. non-activated obstacle avoidance), interface designs, or development stages of 161 the same RR were compared [17², 19, 25-28, 30, 31, 33, 34]; (3) comparisons in a pre/post-test study 162 design with focus on the user experience [34] or the technical functionality [23], assessed before and 163 after/over a series of trials; and (4) comparisons between outcomes of a newly developed robotic 164 monitoring functionality and those of an external criterion measure as a reference measurement $[16^2]$.

165

166 Interventional Studies

Two articles described studies that used an interventional approach, providing training 167 opportunities with the RR [16, 36]. In one study, the subjects' gait performance with the robotic gait 168 assistance system was assessed on six consecutive days [16¹]. However, subjects seemed to use the RR 169 170 only during test procedures and not in their daily routine. Although the ultimate research hypothesis 171 for this 'interventional' approach was lacking, we assumed that the repeated use represented a type of 172 training intervention in order for the subjects to get used to using the RR. In the other study, a four-173 week randomized controlled trial was conducted to evaluate the effects of ambulation training with a 174 RR compared to a traditional rehabilitation therapy method using parallel bars [36]. In this study, 175 assessment methods were used to evaluate the subjects' motor-functional performance after the robot-176 assisted training intervention.

177

178 Statistical Analysis

An inferential statistical analysis of outcomes was included in only three studies [19, 34, 36]. In 25
studies, outcomes were presented using solely descriptive or qualitative data (e.g. frequencies, means,
SDs, and user comments) [12, 16^{1,2}, 17^{1,2}, 18^{1,2,3}, 20-33, 35, 37, 38].

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183 Assessment Methods

184 Assessment measures used in identified studies can be classified into five categories:

185 (1) established clinical performance-based measures assessing subjects' functional ability to 186 perform a requested task by simple quantitative time-, range-, or rating-based outcomes (e.g. gait 187 speed, walking distance, rating score) or by more detailed, qualitative outcomes captured by external 188 technical measures (e.g. step time, double support time); (2) tailored assessment methods in terms of 189 self-designed performance-based measures specifically tailored to specific functionalities of the RR 190 (e.g. guidance system, obstacle avoidance). In addition to simple quantifiable time- or count-based 191 outcomes (e.g. walking time, number of collisions), these assessment methods predominantly used 192 more technique-based and qualitative outcomes (e.g. path deviation, distance to obstacle); (3) 193 assessment methods used to evaluate the subject's physical and physiological demands during the use 194 of the RR; (4) subjective evaluation measures to assess a user's experience with the RR; and (5) 195 technical evaluation measures to assess the technical capability of the RR.

As technical evaluation measures used in nine studies [12, 16^2 , $18^{1,2}$, 20, 22-24, 33], exclusively focused on the technical verification of the RR with limited relevance for the user perspective, we do not further address and discuss these measures in this review.

199

200 Clinical Performance-Based Measures

201 Established clinical performance-based measures were used in three studies [21, 32, 36]. In one of 202 these, the subjects' gait and functional performance with the RR were assessed by the 4-meter walk test (4MWT), a modified version of the TUG, and spatio-temporal gait parameters (i.e. step time, 203 204 double support time) captured by video camera during both tests [21]. Other studies documented the 205 subjects' motor performance by the 6-minute walk test (6mWT), 10-meter walk test (10MWT), and Performance Oriented Mobility Assessment (POMA) [36] or only by the 10MWT [32]. The most 206 207 frequently used outcomes were gait speed [21, 32, 36], completion time [21], or walking distance and 208 rating scores for functional performance (POMA) [36].

In one study, an established screening test for assessing the functional ability of subjects to perform
activities of daily living (ADL) was used (Barthel ADL Index) [36].

211

212 Tailored Assessment Measures

In ten studies, assessment strategies included self-designed performance-based measures 213 specifically tailored to specific robotic functionalities [16^{1,2}, 17², 19, 25-28, 31, 34]. Obstacle 214 avoidance and guidance systems were evaluated while subjects completed walking paths [25, 28] or 215 obstacle courses $[17^2, 31, 34]$, navigation and localization systems while performing navigational tasks 216 [26, 27], and gait assistance systems by analyzing the subject's gait during robot-assisted walking 217 $[16^{1,2}, 19]$. Simple quantifiable outcomes of these tests included number of collisions [26, 31, 34], 218 reorientations [34], navigational mistakes [27] or abnormal gait patterns $[16^{1,2}]$, walking time [34], or 219 220 achievement of task [26]. More specifically tailored, technique-based outcomes, as used in eight studies, comprised of deviations from an optimal path $[17^2, 25, 28, 31]$, distance to obstacles [17, 26], 221 maximum speed and walking distance [26], mean and SD of robot's velocity [19], and gait variability 222 (i.e. SD of gait speed/step length) $[16^{1,2}]$. To obtain such technically advanced outcomes, five studies 223 224 used the data flow created by the technical systems installed on the RR, including laser rangefinders (LRF) [16^{1,2}, 28], a video camera and sonar sensors [17²], or a web camera [31]. In the other three 225 studies, information on the technical measure to capture these outcomes was nonexistent [19, 25, 26]. 226 Out of the studies that determined outcomes with the robot-integrated technical systems, only one 227 228 seemed to process raw data (LRF data) into outcome variables (i.e. path deviation) by using an already 229 established method for robust position estimation of mobile robots in indoor environments ('Monte Carlo localization') [28]. In the other four studies, it remained unclear whether raw data was analyzed 230 by self-designed or potentially established methods $[16^{1,2}, 17^2, 31]$. 231

In two inter-device comparative studies, a bicycle speedometer attached to the conventional device [16] or a LRF placed in the test environment [26] was used to assess technically advanced outcomes such as walking distance or gait variability also when not using the RR. However, a reference, or any information on the psychometric quality of these methods, was missing in both studies.

In four studies including tailored assessment measures, test procedures appear to be nonstandardized $[16^2, 26, 34]$ or have been insufficiently described [28].

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239

Evaluation of Physical and Physiological Demands

240 Four studies assessed subjects' physical and physiological demands with motorized RR during 241 time-based performance-based measures (i.e. navigational trail, 10MWT) [26, 32] or during walking 242 with standardized gait speed [19, 33]. In two studies, the exertion of force applied to steer the RR was 243 measured using the force/torque sensors integrated on the robot's handles [19, 26]. One also reported 244 on forces required to operate a conventional walker, but did not mention the method to capture these 245 forces [26]. The other study additionally evaluated the oxygen consumption (VO₂) and metabolic cost 246 of transport (metabolic cost per unit of mass and distance travelled) during robot-assisted gait using 247 open-circuit respirometry [19]. In the remaining two studies, the muscle activity in the lower 248 extremities was recorded by electromyography (EMG) [32, 33], and one also measured torso 249 kinematics by a tri-axial accelerometer attached to the subject's back [32].

250

251 Subjective Evaluation Measures

252 Nineteen studies included measures to evaluate the subjects' experience with the RR [12, 16¹,17^{1,2},18^{1,3},19, 22-24, 26-30, 34, 35, 37, 38]. However, assessment instruments to perform such 253 subjective evaluations varied widely in methodological quality. Nine studies documented solely non-254 specific comments of non-standardized surveys [16¹, 17², 18¹, 22, 24, 28, 29, 35, 37], three used 255 standardized (dichotomous) questions [27, 30, 38], four used self-designed structured questionnaires, 256 each with different multi-stage rating scales (e.g. 1 to 5, 0 to 100) [12, 17¹, 19, 34], two mentioned the 257 use of questionnaires but did not provide detailed information on contents or a reference $[18^3, 26]$, and 258 259 one presented results of the subjective evaluation by response categories referring to different items but without mentioning the assessment instrument used for this purpose [23]. Most frequently used 260 outcomes of standardized surveys included maneuverability [12, 17¹, 38], safety [12, 30, 38], and 261 262 comfort [12, 19, 34].

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264 Discussion

The aim of this systematic review was to summarize the methodology of evaluation studies of RRs with focus on the user perspective. Identified studies showed large heterogeneity in definitions of potential users, study population, study design, and assessment methods. We found major methodological shortcomings related to insufficient sample descriptions and sample sizes, lack of appropriate, standardized and validated assessment instruments, and lack of statistical analysis of study results. No generic methodology to evaluate RRs could be identified.

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272 User Group Definitions

273 The majority of user group definitions seemed inadequate to guide a technical development of an 274 AAL system. Generic, setting-specific, non-specific impairment-based or disease-oriented definitions 275 do not relate to specific functional impairments of potential users, but cover users with a wide range of 276 different functional abilities and requirements. The effective design of AAL systems in such 277 heterogeneous user groups may be not feasible. The main goal of an AAL system should rather be to 278 overcome or compensate for specific impaired functions. Clear impairment-related definitions are 279 therefore mandatory to specifically tailor AAL developments for specific impairments of users and to 280 ensure that innovative functionalities effectively address a user's needs. When such specific 281 impairment-related definitions are additionally based on standardized and validated assessment 282 methods with established cut-off values, a general comparability of developments and evaluations will 283 be feasible.

Definitions according to impairment levels will in turn allow specifications such as risk stratification of potential users. With this, the user group will be further classified opening up the option to exclude persons with no or minor impairment, with no need for assistive devices, or with advanced impairment or unacceptable risk exposure when using the device (triage). Another specification may focus on the main function of the specific device. For example, when an AAL system such as a RR basically supports gait performance, a specific definition based on standardized and validated gait assessment (e.g. 10MWT) will be superior compared to less specific definitions such as general functional scores (e.g. Barthel ADL Index).

As the user group of RRs may be old and multi-morbid persons, also highly prevalent ageassociated impairments might be included in the definitions, depending on the specific functionalities or complexity of devices (e.g. inclusion of cognitive impairment with respect to navigation functions in disoriented persons).

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297 Study Samples

Overall, sample sizes seemed rather limited to give a consistent picture of the user perspective. Surprisingly, the statistical analysis of documented data was not in the focus of studies as only a very limited number included such analyses (3 of 28) and none of these presented a sample size calculation as a prerequisite of statistical analysis.

A remarkable number of studies (10 of 28) evaluated RRs in persons who were not covered by the predefined user group, considerably limiting the user perspective of these studies. Study results with inadequate, convenient, or insufficiently described samples may not suffice to allow conclusions for persons with specific impairments which may represent the potential users of the RR. To ensure that RRs meet a user's needs and requirements and become successful in the market, it seems mandatory to involve the intended users at all stages of the design and evaluation process of such assistive robotic technologies [39-41].

309

310 **Design of Studies**

311 Observational Studies

312 The most heterogeneous group of studies covered observational studies that used solely descriptive 313 data presentations without providing any reference or comparative values. Findings and conclusions of 314 these studies were thus mainly based on the authors' subjective perception and appraisal. However, 315 when using standardized and validated outcome measures with well-established cut-off values or other 316 assistive mobility devices for comparison, such observations lose their merely subjective and study-317 specific nature and enable the objective appraisal of outcomes related to other studies or the 318 documentation of an added value of the RR compared to other devices. From a user as well as a 319 technical perspective, observational studies that descriptively presented non-classifiable or non-320 comparable outcomes therefore seem to have limited value.

321

322 *Comparative Studies*

The documentation and perception of an added value of the RR is of utmost importance for potential users. Innovative high-tech developments may be fascinating and mandatory for engineering research; however, they may also lead to rather complicated devices for everyday use, not easy to maneuver, too complex to operate, or too expensive to afford. A comparison of RRs with established, low-tech devices ('inter-device comparative study design') may therefore be useful to demonstrate to users the benefit of RR usage.

Comparisons may also be used for the evaluation of single functionalities to document the effect of a specified functionality (e.g. activated guidance system) or the progress of a new development stage. Such an 'intra-device comparative' study design allows a tailored assessment of the subjects' functional performances, physical and physiological demands, and user experience in specific assistance levels or development stages of the RR.

Frail, older persons may initially be intimidated by the robot's appearance in early stages of development (e.g. without casing, exposed hardware) which may in turn result in a more negative user perception before actually having used the RR. Subjective user evaluations, in a pre/ post-test study design, provide the opportunity to assess the subjects' initial impressions of the RR and whether there are potentially negative prejudices, which may, however, be overcome after actual use of the RR.

Independent of different types of comparative studies, such a study design should definitely includea statistical analysis to compare results which was however seldom used in the identified studies.

341

342 Interventional Studies

343 An interventional study design represents a new aspect in evaluation studies with strong focus on 344 the user perspective. Newly developed RRs may not necessarily meet a user's acceptance or provide 345 usability and efficiency when using them for the first time. Insufficient training opportunities or 346 instruction prior to assessment measures may jeopardize study outcomes [42]. An adequate practice 347 time therefore seems mandatory to prevent initial problems in operating the RR, and may further 348 increase the impact on outcomes. Particularly when comparing RRs with a subject's own conventional 349 assistive devices, brief instructions may not be sufficient, as subjects are already much more familiar 350 and better trained with their own devices.

351 Overall, we identified a lack of studies investigating usability of RRs in natural environments with 352 adequate long-term evaluation of habitual use. The development and evaluation of RRs seemed to 353 occur rather in engineering laboratories than in clinical settings, as already reported for other robotic 354 assistance systems (e.g. service robots, robotic exoskeleton) [43]. This may be explained by the fact 355 that most of the identified studies evaluated research prototypes in rather early development stages, 356 not yet ready for market launch. In such stages, it is important to manipulate specific variables of a 357 prototype in order to investigate their effects precisely and to optimize technical functionalities 358 accordingly [41]. Since laboratory evaluations also require less time and provide highly standardized 359 conditions, a restricted experimental study design may have been favored. However, for the ultimate 360 goal of RRs to assist mobility of impaired persons in daily life, tests for habitual use seem to be 361 mandatory documenting risk, experience-based perception of use, and quality of life with high 362 relevance for users as well as caregivers.

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364 Assessment Methods

365 Clinical Performance-Based Measures

Internationally well-established, clinical performance-based measures allow a worldwide 366 367 comparability of results, but may be insufficient to cover the particular added value of specific robotic 368 functionalities (e.g. obstacle avoidance, navigation assistance) as the outcome variables do not 369 necessarily refer to the subjects' abilities potentially affected by the RR [42]. In addition, clinical 370 assessment methods may be limited by subjective rating (POMA) or limited with respect to less 371 detailed, unidimensional outcomes such as gait speed (4MWT, 10MWT) or task completion time 372 (TUG). Augmenting such measures with technical assessment systems (e.g. video analysis system) 373 allows a multidimensional analysis of the subjects' gait, including outcomes related to insecure gait or 374 postural (in-)stability (e.g. width of base of support, double vs. single limb support) and reduction of 375 falling risk as a main target of RRs.

Even established and validated assessment methods may have their limitations when inadequately used. Outcomes such as gait speed (4MWT) and task completion time (TUG) may be inappropriate when comparing a non-motorized, conventional device with a motorized RR with limited maximum speed. In such comparisons, a superior outcome for the low-tech device seems almost mandatory and may indicate an insufficient selection of a study outcome. The use of ADL scales (e.g. Barthel ADL Index) to evaluate effects of a robot-assisted ambulation training appears also inappropriate, since they include, if any, only very few sub-items targeting the subject's walking ability.

Another potential methodological pitfall may be related to performance-based outcome variables with ambiguous consequences: a motorized RR will improve gait speed in less impaired persons without substantial risk. However, improved performance may be traded off by a substantially higher risk of falling in more impaired persons.

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388 Tailored Assessment Methods

389 The quality of an assessment strategy substantially depends on the appropriateness of methods with 390 focus on the newly developed functionalities to document the added value of RRs. Clinical 391 performance-based measures may be attractive because of their well-established psychometric 392 properties; however, they have been developed for clinical purpose and may not cover new 393 functionalities in innovative assistive technologies [42]. An assessment strategy specifically tailored to 394 the specific functionality to be evaluated may help to achieve this goal. In RR, depending on the 395 functionalities installed, a huge data flow created by the robot-integrated sensing technique already 396 exists to control motoror cognitive assistance systems. Using this data flow for assessment purposes 397 may allow highly qualitative and quantitative tailored assessments exactly tuned to the newly 398 developed functionality in order to document the added value of the RR. For example, when focusing 399 on functionalities providing navigational assistance, the data flow from laser sensors, which is used to 400 feed back the position of the RR, could be processed into a superior assessment of walking trajectories during a navigational task. When using such data for assessment purpose, it seems mandatory to 401 402 examine or to provide sufficient information on the psychometric qualities of the robot-integrated 403 sensor technique and the analysis method used to process raw data into the outcome variables. 404 However, it appeared that only one study used an already established method for this approach [28]. 405 Furthermore, to ensure reliable, reproducible, and comparable outcomes, the test procedure of tailored 406 assessment measures has to be also clearly standardized.

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408

Evaluation of Physical and Physiological Demands

409 Measures such as EMG, respirometry, accelerometry, or measurements of applied steering forces 410 to the RR allow a detailed insight into relevant physical and physiological effects on objective 411 parameters, which may be indicators for the subject's individual physical exertion (e.g. VO₂, muscle 412 activity). However, some of these rather laborious measures (e.g. EMG, respirometry) seem less 413 amenable for old and multi-morbid persons and may have therefore been used predominantly in 414 studies including only young, healthy adults [19, 33]. To prevent overtaxing by test conditions, 415 alternative methods to evaluate physical exertion are available which may increase amenability by 416 standardized and validated subjective rating (e.g. [44]).

417

418 Subjective Evaluation Measures

419 In studies including subjective evaluation measures, a wide range of methods (e.g. non-specific 420 comments, self-designed questionnaires) related to a variety of different aspects of the subject's 421 experience with the RR was used which may considerably limit the comparability of outcomes. The 422 overall lack of already established, validated questionnaires for the subjective evaluation of assistive 423 technology (e.g. [45-47]) might be due to two reasons: (1) established questionnaires have been 424 developed for a generic evaluation of a wide range of assistive technology devices but may be limited 425 for evaluating specific functionalities of individual devices [45]; (2) some questionnaire items may 426 also be inappropriate to evaluate prototypes after a short-term experiment in a restricted test scenario, 427 covering aspects such as quality of life, usability in daily routine, durability, or services [45-47] whose 428 assessment may only be feasible after habitual use of the devices over an extended period of time. 429 However, the subjective evaluation measures used in the identified studies rather targeted the subject's 430 actual experience directly after using the RR. This may explain the use of self-developed 431 questionnaires including items already assessable after short-term use in an artificial setting (e.g. 432 maneuverability, safety, ease of use). However, only when these questionnaires have been validated before application and internationally established cut-off values are available, such assessmentinstruments guarantee high psychometric quality and allow comparability of study results [48].

435

436 Limitations

437 Only information available in the articles was evaluated in this review, although the authors may have used additional or more detailed methodology, not stated in articles. The fact that the evaluation 438 of AAL prototypes may require elaborate and costly ethical application and study procedures 439 440 ('Medical Product Act') may have prevented RRs to be tested in comprehensive studies with adequate 441 sample sizes and the target user group as well as in natural environments with adequate long-term 442 evaluation of habitual use. The role of clinical partners in AAL research projects may offer 443 opportunities to solve such problems. Clinical partners may be able to provide specific impairment-444 based user group definitions, to recruit a satisfactory number of potentially adequate subjects, and to 445 investigate the habitual use of AAL systems in natural environments.

446

447 **Conclusions**

448 Apart from the heterogeneity, methodological deficits in most of the identified studies became 449 apparent. Recommendations for future evaluation studies include: (1) clear definition of target user 450 group by valid, specific impairment-based criteria; (2) adequate selection of subjects with predefined 451 inclusion criteria representative for potential users; (3) inclusion of other assistive mobility devices for 452 comparison; (4) inclusion of the habitual use of advanced prototypes in evaluation rather than mere 453 short-term, restricted, experimental test scenarios for single functionalities of prototypes not finalized 454 for use in the target user group; (5) selection of established, standardized, and validated assessment 455 methods; (6) implementation of a specifically tailored assessment strategy, focusing on specific 456 functionalities of the RR, and (7) statistical analysis of study results. These recommendations, given 457 for RRs, may also apply in general for the development and evaluation of AAL systems with focus on 458 the user perspective.

459

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467

468 **Conflict of Interest**

469 The authors have no conflict of interests to declare.

470

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Assistive mobility device	Robotic functionality	Gait/mobility support	Evaluation measure
#1 'robotics'[Mesh]	#14 'electric power supplies'[Mesh]	#23 'gait'[Mesh]	#32 'evaluation studies as topic'[Mesh]
#2 'walkers'[Mesh]	#15 robot*[tiab]	#24 'Walking'[Mesh]	#33 'Technology Assessment, Biomedical'[Mesh]
#3 'self-help devices'[Mesh]	#16 smart[tiab]	#25 'Dependent Ambulation'[Mesh]	#34 evaluat*[tiab]
#4 'biomedical technology'[Mesh]	#17 intelligent[tiab]	#26 gait[tiab]	#35 assess*[tiab]
#5 robot*[tiab]	#18 power*[tiab]	#27 walk*[tiab]	#36 measur*[tiab]
#6 rollator*[tiab]	#19 electric[tiab]	#28 ambulant*[tiab]	#37 trial*[tiab]
#7 mobile platform*[tiab]	#20 motorized[tiab]	#29 mobility[tiab]	#38 experiment*[tiab]
#8 mobility aid*[tiab]	#21 motorised[tiab]	#30 OR (#23-#29)	#39 test*[tiab]
#9 mobility device*[tiab]	#22 OR (#14-#22)	#31 (#13 AND #22 AND #30)	#40 clinical[tiab]
#10 assistive device*[tiab]	#23 (#13 AND #22)		#41 OR (#32-#40)
#11 assistive system*[tiab]			#42 (#13 AND #22 AND #30 AND 41)
#12 walking aid*[tiab]			
#13 OR (#1-#12)			

Table 1. Overview of the search term used in PubMed

Name of device Authors [Ref. No.]	User group definition	Study sample	Study object	Study design	Assessment methods Type: outcome measurement: outcome variable
Context-aware PD patients Assisted Interactive RObotic Walker	PD patients	patients $\frac{Study \ 1}{n = 6}$ $Age: n/a$	UP	IV: repeated assessment on six consecutive days	TAM: gait analysis on straight walking path ^b ; CAIROW gait analysis system (based on LRF) ^b : SD of gait speed/step length; expert rating of gait ^d : number of abnormal gait patterns (festinating gait, freezing of gait)
(CAIROW) Mou et al. 2012 $[16^{1,2}]$		PD patients of senior care unit; mHY, stage range 1.5-3	UE	OB	SEM: user comments ^c after gait analysis
	Study 2 n = 7 (F = n/a) Mean age: 86 yrs PD patients of senior care unit;	UP	Inter-DC: walking with CAIROW vs. normal walking (with own/ with- out assistive device)	TAM: gait analysis on walking path with obstacles, people randomly passing by, up- and down-going slopes, short section for backward walking ^d ; CAIROW gait analysis system ^b or LRF ^c placed in test environment when normal walking: SD of gait speed/step length; expert rating of gait ^d : number of abnormal gait patterns	
		mHY, stage range 1-3	TC (gait analysis system)	EC: gait analysis system vs. expert rating	TEM (see original article for details)
Care-O-bot II Graf 2009 [26]	Elderly people in home environment	n = 6 (F = 5) Age range: 86-92 yrs Inhabitants of an old people's residence using mobility aids in daily life	UP, PD	Inter-/intra-DC: target mode (robot-determined motion control) vs. direct control mode (user-deter- mined motion control) vs. conventional walker	TAM: navigation trail in old people's residence with a ramp, tables, and people randomly passing by ^d ; robot's guidance system ^c , bicycle speedometer ^c mounted on conventional walker: walking time, number of collisions, maximum speed, walking distance, distance to obstacle PHY: force/torque sensors ^c in robot's handles, force measurement when using conventional walker not reported ^d : pushing force
			UP	OB	TAM: navigation trail in old people's residence with transition between ground floor and 1^{st} floor, a ramp, tables, people randomly passing by ^d : achievement of target
			UE	OB	SEM: questionnaire ^b after navigation trail: n/a
Chugo group walker Chugo et al. 2009 [30]	Elderly people in need for nursing in daily routine	$\label{eq:response} \begin{array}{l} n=7 \; (F=n/a) \\ Age: \geq 67 \; yrs \\ People \; in \; need \; of \; long-term \; care \; at \\ level \; I \; or \; II \; in \; Japanese \; Long-term \\ Insurance \; System \end{array}$	UE	Inter-/intra-DC: STS transfer without assistance vs. with previous/novel STS assistance system	SEM: questionnaire ^b after STS transfer : ease of standing up, fear of falling (1= inferior, $3 = \text{same}$, $5 = \text{better}$ feeling compared to STS transfer without assistance)
CO-Operative Locomotion Aide (COOL-Aide) Wasson et al. 2008 [22]	Elderly people n = 12 (F = Mean age () Healthy sub with disord (cerebral pa	Iderly people $n = 12$ (F = 2)Mean age (SD): 36.8 (18.1) yrsHealthy subjects (n = 8), subjectswith disorders affecting mobility(cerebral palsy, familial torsiondystonia) (n = 8)note: (1) total sample, (2) - (5)subsample: only healthy subjects	TC (guidance, user intent detection and obstacle avoidance system)	OB	TEM (see original article for details)
			TC (obstacle avoidance system with vs. without stability preservation)	Intra-DC: standard vs. stability-preserved obstacle avoidance	TEM (see original article for details)
			UE	OB	SEM: user comments ^d after performing a set of short obstacle courses
Gait Rehabilitation	Disabled or elderly	n = 2 (F = 0) Mean age (SD): 28 5 (2.1) yrs	TC (guidance system)	OB	TEM (see original article for details)
(GRSR) Jang et al. 2008 [33]	problems or paralysis; weighing up to 75 kg	Ordinary adult males	PD	Intra-DC: 40/20 % body weight support vs. full body weight	PHY: EMG ^a during straight walking with standardized gait speed of 0.2 m/s: muscle activity of lower extremities (EMG signal) (quadriceps, hamstrings, gastrocnemius, tibialis anterior)

Table 2. Study cl	haracteristics and	l assessment me	ethods of the	28 studies inclu	ded in this systematic	review

Table 2. (continued)

Name of device Authors [Ref. No.]	User group definition	Study sample	Study object	Study design	Assessment methods Type: outcome measurement: outcome variable
GuidoFrail elderly peopleRentschler et al. 2008with visual[34]impairment	n = 17 (F = n/a) Mean age (SD): 85.3 (7.0) yrs Residents of a supportive living facility/nursing home with visual impairment due to macular degeneration, cataract, glaucoma or other reasons; mean time (SD) since onset of visual impairment: 20.4 (13.0) yrs; ambulatory (\geq 20 min within 90 min period) with	UP	Inter-/intra-DC: Guido vs. conventional assistive mobility device or normal walking (with own/ no assistive device); automatic (user- determined motion control) vs. manual mode (shared user-robot motion control)	TAM: obstacle course with randomly placed obstacles before each trial ^d : walking time, number of obstacle/wall collisions, number of reorientations	
		limited assistance	UE	PPC: before and after 3 trials	SEM: Subjective Mobility Questionnaire ^{<i>b</i>} after obstacle course: appearance, ease of use, usefulness in living environment, embarrassment ($1 = best score; 5 = worst score$)
Hitachi walker Tamura et al. 2001 [32]	Elderly people who have difficulty walking	$ \begin{array}{l} n=6 \; (F=n/a) \\ \mbox{Mean age (SD): 82 (7.9) yrs} \\ \mbox{Subjects ambulatory with} \\ \mbox{supervision } (n=4), \mbox{subjects in} \\ \mbox{need for walking assistance } (n=2) \end{array} $	UP PD	Inter-DC: Hitachi vs. caster vs. conventional walker; robot vs. parallel bars	CPM:10MWT ^a : gait speed PHY: EMG ^c , tri-axial accelerometer ^c during non-standardized gait speed (10MWT): muscle activity (EMG signal), trunk acceleration
HUST walking-aid robotElderly or disabled peopleXu et al. 2013 [23]	Elderly or disabled people	disabled n = 3 (F = n/a) Age: n/a Volunteering subjects with/ without experience using robot;	TC (motion control system)	PPC: autonomous learning process of HUST in motion behavior over a series of trials	TEM (see original article for details)
	joint to imitate lower limb disorders	UE	OB	SEM: subjective evaluation after completing a series of obstacle courses, assessment measure not reported ^{d} : flexibility, comfort, maneuverability, obstacle avoidance	
i-Go Ko et al. 2014 [24]	Elderly people	n = 3 (F = n/a) Age: "in their twenties"	TC (guidance system) UE	OB	TEM (see original article for details) SEM: user comments ^d after completing an S-shaped walking path
Intelligent Mobility Platform (IMP) Glover 2003 [29]	Older adults (primarily without major visual or cognitive impairment)	n = 6 (F = n/a) Age: n/a Residents of a care facility with/ without need for walker	UE	OB	SEM : user comments ^{d} after presentation and informal testing of the robot
iWalker Persons with stree early- to mid-sta [27] AD, traumatic brinjury, macular degeneration, cataracts, visual impairment; primarily in nurs and assisted livin homes	Persons with stroke, early- to mid-stage AD, traumatic brain injury, macular	with stroke, $n = 4$ (F = n/a) age: n/a umatic brain nacular service currently using cane, ation, walker or bot, with history of way s, visual finding problems; MMSE, mean sent; score (SD): 26 (3.6) y in nursing sted living	UP	Inter-DC: iWalker vs. conventional device (cane/walker) accompanied by researcher	TAM: several navigation trails ^{b} : walking time, number of navigational mistakes
	degeneration, cataracts, visual impairment; primarily in nursing and assisted living homes		UE	Intra-DC: map-based (+ auditory cues) vs. text-and- arrow-based (+ auditory cues) user interface design	SEM: dichotomous question ^b : choice of user interface; user comments ^d

Table 2. (continued)

Name of device Authors [Ref. No.]	User group definition	Study sample	Study object	Study design	Assessment methods Type: outcome measurement: outcome variable
i-Walker (EU) Annicchiarico 2012 [36]	Post-stroke patients with hemiparesis	$\label{eq:states} \begin{array}{l} n = 20 \ (F = 11) \\ \mbox{Mean age: 59.9 yrs} \\ \mbox{Acute hemiparetic stroke patients} \\ (event < 1 \ yrs) \ receiving \\ rehabilitation treatment; \ \mbox{MMSE} \\ \mbox{score} \geq 20; \ \mbox{CNS upper \& lower} \\ \mbox{limb} > 0 \end{array}$	UP	IV (RCT): robot-assisted ambulatory training (EG) vs. in parallel bars (CG) (4 weeks, 5x a week)	CPM: POMA ^{<i>a</i>} : total score; 6mWT ^{<i>a</i>} : walking distance; 10MWT ^{<i>a</i>} : gait speed ADL screening: Barthel ADL Index ^{<i>a</i>} : score
i-Walker (Japan) Kikuchi et al. 2010 [31]	Patients with imbalanced motor/sensory functions (e.g. hemiplegic patients), difficulties in smooth walking	n = 6 (F = 2) Mean age (SD): 88.7 (6.1) yrs Residents of elder care facility with wheelchair due to loss of vision/muscle strength which occasionally train walking with forearm caster walker; chronic disease: stroke, dementia, muscle atrophy, high blood pressure, heart failure, AD, cataract, PD	UP	Intra-DC: passive vs. active robot motion control system	TAM: walking path with obstacles ^{<i>b</i>} , robot-integrated web camera ^{<i>c</i>} : deviations from a path marked on the floor, number of collisions
JAIST Active Robotic Walker (JARoW) Lee et al. 2014 [38]	Elderly people with certain level of ambulatory capability (FAC score 4-5)	n = 5 (F = 4) Age range: 75-84 yrs Subjects using traditional walkers in daily routine	UE	Inter-DC: JARoW vs. conventional walker	SEM: questionnaire ^b after walking around for 10 min: ease of walking, safety, maneuverability, suggestions for improvements
MOBIL walking & lifting aid Bühler et al. 2001 [18 ¹]	Frail, elderly and walking disabled people	$\label{eq:study_l} \begin{array}{l} \underline{Study \ l} \\ n \geq 2 \ (F = n/a) \\ Age: n/a \\ Selected users, technical and \\ rehabilitation experts \end{array}$	TC (overall system functionality) UE	OB	TEM (see original article for details) SEM: user/expert ratings, comments and interviews ^d
MOBIL test bed [18 ²]	Frail, elderly and walking disabled people	$\label{eq:study_2} \begin{array}{l} \underline{Study \ 2} \\ n \geq 2 \ (F = n/a) \\ Age: n/a \\ Rehabilitation engineers, walking \\ impaired persons \end{array}$	TC (overall system functionality)	OB	TEM (see original article for details)
MOBIL walking & lifting aid, MOBIL test bed [18 ³]	Frail, elderly and walking disabled people	$\label{eq:study_3} \begin{split} \underline{Study \ 3} & n \geq 2 \ (F = n/a) \\ Age: n/a \\ Community-dwelling people, \\ institutionalized elderly disabled \\ people, \ care \ staff \end{split}$	UE	OB	SEM: questionnaire ^b after demonstration, video presentations, practical trials: n/a

I unic 2. (continueu)	Table 2.	(continu	ied)
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Name of device Authors [Ref. No.]	User group definition	Study sample	Study object	Study design	Assessment methods Type: outcome measurement: outcome variable
Nomad XR 4000 Morris et al. 2003 [28]	Frail older people with cognitive impairment	n = 4 (F = n/a) Age: n/a Residents of a retirement facility	UP	Intra-DC: passive (no navigational assistance) vs. active (with navigational assistance) vs. forced mode (full robot motion control)	TAM: navigational trail ^d ; robot's navigation system (based on LRF, 'Monte Carlo localization') ^a : deviation from optimal path
			UE	OB	SEM: user comments after navigational trails ^d
Personal Aid for Mobility and Monitoring (PAMM SmartWalker) Yu et al. 2003 [17 ^{1,2}]Independently living or institutionalized elderly people with mobility difficulties due to physical frailty and/or disorientation due to age and sickness		UE	OB	SEM: questionnaire ^b after free driving at facility: ease of control, going straight, turning, heaviness, support, satisfaction $(1 = \text{worst score}, 5 = \text{best score})$	
	frailty and/or disorientation due to age and sickness	Study 2 n = 8 (F = 5) Age range: 84-95 yrs Elderly residents of assisted living facility with need for walkers	UP	Intra-DC: full robot motion control vs. adaptive shared user-robot motion control vs. without any motion control	TAM: wall-limited walking path through assisted living facility ^d ; robot's vision-based localization system (based on charged-coupled device camera) ^b : deviations from robot-generated, pre-planned path, distance to wall
			UE	OB	SEM: user comments ^d
Robotic Mobilityn/aPlatform (RMP)Grondin & Qinggou2013 [19]19	n/a	n = 10 (F = 5) Mean age (SD): 24.6 (3.0) Subjects without previous/current gait-related injuries and without experience in using rollators or robotic walkers	UP, PD	Intra-DC: novel vs. previous motion control system	TAM: walking with targeted velocity of 1 m/s through a circular path in low-traffic hallways ^b ; technical outcome measurement not reported ^d : mean and SD of robot velocity; PHY: force/torque sensor ^a under robot's left handle: pushing force
			PD	Inter-/intra-DC: novel vs. previous motion control system vs. conventional rollator vs. no assistive device	PHY: walking with targeted velocity of 1 m/s through the circular path ^c (use of a Hall effect sensor mounted on the conventional rollator to display target velocity); respirometry ^{<i>a</i>} : metabolic cost of transport, oxygen consumption
			UE	Intra-DC: novel vs. previous motion control system	SEM: questionnaire ^b : comfort, intuition, speed control, exertion, overall experience (0 = worst score, 5 = best score)
robuWALKER Rumeau et al. 2012 [21]	elderly people	$\begin{array}{l} n=8\ (F=5)\\ Mean age\ (SD):\ 82.6\ (8.7)\ yrs\\ Healthy\ elderly\ (n=4):\ 4MWT < \\ 4s,\ TUG < 13s,\ MMSE\ score \geq 26;\\ elderly\ patients\ with\ motor\ & \\ cognitive\ impairment\ (n=4):\\ 4MWT > 4s,\ TUG > 13s,\ MMSE\\ mean\ score\ (SD):\ 20\ (3.5);\ all\\ subjects\ without\ experience\ in\\ using\ walking\ frames\end{array}$	UP	Inter-DC: robuWalker vs. conventional walker	CPM: 4MWT ^a : gait speed, modified TUG ^a : completion time; gait analysis by video recordings ^c during 4MWT and TUG: step time, double support time

 Table 2. (continued)

Name of device Authors [Ref. No.]	User group definition	Study sample	Study object	Study design	Assessment methods Type: outcome measurement: outcome variable
Robotic Travel Aid (RoTA) Mori et al. [35]	visually impaired community-dwelling people, hospital patients, or residents of senior homes loss of ability to walk with mobility aids for the blinds	n > 60 (F = n/a) Age: n/a Blind and weak-sighted elderly people	UE	OB	SEM: user comments ^d after walking course
RT Walker Taghvaei et al. 2010 [20]	n/a	n = 2 (F = n/a) Age: n/a	TC (motion control system)	OB	TEM (see original article for details)
SIMBIOSIS Walker Frizera-Neto et al. SCI patien using whe	SCI patients mainly using wheelchair,	SCI patients mainly ising wheelchair, wut usually able to valk for short $n = 8 (F = n/a)$ Age: n/a Subjects with preserved cognitive functions; ability to (1) maintain standing position, (2) walk 10 m without assistance of another person and with or without support of a mobility aid, and (3) to grasp; WISCI II, mean score (SD): 15.9 (2.9)	TC (user intent detection system)	OB	TEM (see original article for details)
2011 [12]	but usually able to walk for short periods of time with assistance of device, WISCI II = 16		UE	OB	SEM: questionnaire ^b after completing U-shaped walking path: maneuverability, safety, posture & comfort ($0 =$ worst score, $100 =$ best score)
Smart Mobile Walker (SMW) Lee et al. 2012 [37]	elderly people, people with hemiplegia, people with incomplete SCI	n ≥ 2 (F = n/a) Age: n/a Stroke patients, SCI patients, clinical experts	UE	OB	SEM: user comments/interviews ^d after demonstrations
Walking Helper Hirata et al. 2005 [25]	elderly people, disabled people	n = 8 (F = n/a) Age: n/a	UP	Intra-DC: novel vs. traditional motion control system	TAM: following S-shaped walking path ^{b} (marked on the floor); technical outcome measurement not reported ^{d} : deviation from path marked on the floor

Abbreviations: PD = Parkinson's disease; F = females; n/a = not available; mHY = modified Hoehn and Yahr Scale; UP = User performance; UE = User experience; IV = interventional; OB = observational; TAM = tailored assessment measure; LRF = laser rangefinder; SD = standard deviation; SEM = subjective evaluation measure; TC = technical capability; inter-DC = inter-device comparative; EC = comparison with external criterion measure; TEM = technical evaluation measure; PD = physical/physiological demands; intra-DC = intra-device comparative; PHY = evaluation of physical or physiological demands; STS = sit-to-stand; EMG = electromyography; PPC = pretest-posttest comparative; <math>CPM = clinical performance-based measure; 10MWT = 10-meter walk test; MMSE = Mini-Mental State Examination; CNS = Canadian Neurological Scale; RCT = randomized controlled intervention trial; EG = experimental group; CG = control group; POMA = Performance Oriented Mobility Assessment; 6mWT = 6-minute walk test; ADL = activities of daily living; AD = Alzheimer's disease; FAC = Functional Ambulation Classification; TUG = Timed Up and Go; 4MWT = 4-meter walk test; WISCI = Walking Index for Spinal Cord Injury.

^a established, standardized and validated assessment test or outcome measurement.

^b standardized, but not validated test procedure or outcome measurement.

^c potentially an established outcome measurement, but no reference given.

^{*d*} non-standardized or unclear test procedure or outcome measurement.