ABSTRACT

Impressive advances in inhalation therapy for patients with asthma and chronic obstructive pulmonary disease (COPD) have occurred in recent years. However, important gaps in care remain, particularly relating to poor adherence to inhaled therapies. Digital inhaler health platforms which incorporate digital inhalers to monitor time and date of dosing are an effective disease and medication management tool, promoting collaborative care between clinicians and patients, and providing more in-depth understanding of actual inhaler use. With advances in technology, nearly all inhalers can be digitalized with add-on or embedded sensors to record and transmit data quantitating inhaler actuations, and some have additional capabilities to evaluate inhaler technique. In addition to providing an objective and readily available measure of adherence, they allow patients to interact with the device directly or through their self-management smartphone application such as via alerts and recording of health status. Clinicians can access these data remotely and...

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during patient encounters, to better inform them about disease status and medication adherence and inhaler technique. The ability for remote patient monitoring is accelerating interest in and the use of these devices in clinical practice and research settings. More than 20 clinical studies of digital inhalers in asthma or COPD collectively show improvement in medication adherence, exacerbation risk, and patient outcomes with digital inhalers. These studies support previous findings about patient inhaler use and behaviors, but with greater granularity, and reveal some new findings about patient medication-taking behaviors. Digital devices that record inspiratory flows with inhaler use can guide proper inhaler technique and may prove to be a clinically useful lung function measure. Adoption of digital inhalers into practice is still early, and additional research is needed to determine patient and clinician acceptability, the appropriate place of these devices in the therapeutic regimen, and their cost effectiveness.

**Keywords:** Adherence monitoring; Asthma; COPD; Digital health; Electronic devices; Exacerbation; Inhalation device; Medication adherence; Peak inspiratory flow; Remote patient monitoring

### Key Summary Points

1. Digital inhaler devices have existed for over two decades but are only beginning to emerge as an important component of e-health for asthma and COPD management.

2. These devices gather data on adherence, and can be linked with information on symptoms, physiological measures, and environmental conditions to allow personalized decisions about asthma and COPD management.

3. Key roles of these devices include: characterizing and improving inhaler adherence and use; reducing exacerbations; improving inhaler technique and pulmonary function; and informing costly and potentially risky interventions.

4. There are opportunities to improve patient medication adherence and outcomes by using the data from digital inhalers and associated platforms to enhance clinical decision-making, improve adherence, and guide clinical care.

5. More data are needed regarding patient and end-user acceptability, cost-effectiveness, and effect on inhaler technique.

### DIGITAL FEATURES

This article is published with digital features, including a video abstract, to facilitate understanding of the article. To view digital features for this article go to https://doi.org/10.6084/m9.figshare.14816532.

### INTRODUCTION

Aerosol therapies are essential for treatment of obstructive lung diseases, which affect nearly 15% of the global population [1]. Inhaled therapies are some of the most complex therapeutic modalities for patients with chronic diseases to self-administer. Optimal inhaled therapy requires specific breathing maneuvers that are coordinated with the release of aerosol from the inhaler. Moreover, adherence to treatment regimens is often suboptimal [2–4]. Poor adherence to inhaled therapies in both asthma and chronic obstructive pulmonary disease (COPD) is linked with poorer outcomes and health status, increased exacerbations and hospitalizations, and death [5–8]. These problems persist
Despite decades of efforts to address adherence and inhaler technique [9].

Despite their availability in the last two decades, digital inhaler devices, also known as electronic adherence monitors, electronic monitoring devices, electronic medication sensors, or digital inhalers, are only now emerging as an important component of e-health for the management of patients with asthma and COPD [10]. For the purposes of this paper, these devices will collectively be referred to as “digital inhalers.” Broadly speaking, for obstructive lung diseases, e-health includes these digital inhaler devices and physiological monitors, as well as text messaging and clinical platforms using web and mobile applications. Data gathered using digital inhalers on patients’ medication adherence, particularly when supported with information on symptoms, physiological measures, and environmental conditions, can provide a more complete basis for timely and personalized decisions about management [10–12]. The potential to improve patient care with digital inhalers has been significantly enhanced by incorporating newer technology into these devices such as the ability to record location data of the inhaler and to provide real-time feedback on the inhalation technique [10, 13–15].

In this article, we review commercially available digital inhalers and digital sensors (Propeller Health, Teva Digihaler®, Adherium Hailie®, and Amiko Respiro®) and their supporting digital health platforms in detail. We also provide perspectives on the INCA device that has undergone extensive investigation in Ireland and the United Kingdom (UK) but is not currently marketed. Digital inhaler devices by Cognita Labs (California) and others are not discussed due to their current limited availability. We discuss potential clinical applications and review studies that have described the use of digital inhalers in patients with asthma or COPD.

METHODS

We searched Ovid MEDLINE, EMBASE, Cross-Ref, and Google Scholar for English-language publications of randomized controlled trials (RCTs), systematic reviews, and guidelines with Medical Subject Headings for medication adherence, measurement, electronics, and lung disease, and terms related to “electronic monitoring devices,” “digital inhalers,” “electronic medication monitors,” and “adherence monitoring” from January 1, 2010, to April 1, 2021, to cover the last decade of research. Studies were included if digital inhalers (electronic monitoring devices [EMDs] and related mobile applications) were used as part of an adherence intervention and had the primary aim of improving medication adherence, and were conducted in patients with asthma or COPD. Studies were excluded if they did not report results using the Consolidated Standards of Reporting Trials (CONSORT) criteria or similar standardized reporting methods, or had follow-up periods less than 3 months. Studies were also identified through a manual search of the reference lists of the literature and based on expert advice from clinicians and researchers with experience working with digital inhalers.

Compliance with Ethics Guidelines

This article is based on previously conducted studies and does not contain any new studies with human participants or animals performed by any of the authors.

Digital Inhaler Health Platforms and Devices

Digital inhaler health platforms comprise an inhaler digitalized with electromechanical sensor(s) and associated microelectronics to detect the time/date of inhaler actuations. Other features included in some devices are a connected smartphone, a dedicated mobile application that receives data and interacts with the patient, a cloud server, and a portal for the clinician to review data transmitted to their dashboard (Fig. 1). These devices are for single person use, and sensors can be used for both rescue and controller inhalers. All digital inhaler platforms are compatible with iOS and Android smartphones and their data are encrypted.
For platforms which track location data, the location of the “inhaler” is based on the location of the patient’s smartphone, and it can be linked to environmental reports. At present, the Global Positioning System (GPS) cannot be used to locate the inhaler, as data are drawn from the smartphone rather than location of the inhaler itself [16], which may limit the accuracy of linking location data with inhaler use and/or tracking environmental exposure to triggers and association with asthma exacerbations.

The most common commercially available digital inhaler devices for monitoring inhaler use are (1) Propeller Health—global, (2) Teva Digihaler®—USA, (3) Amiko Respiro®—Italy, Germany, Portugal, Netherlands, and (4) Adherium Hailie®—global. The INCA device (Ireland and the UK) is not currently marketed; however, published research on this device has added significantly to our understanding of this technology, and it is included in this review. Table 1 provides details about these devices and medications that are compatible for use with each inhaler device.

The electronic sensors are either attached (Propeller Health, Hailie®, Respiro®, INCA) [17] or embedded in the inhaler (Teva Digihaler®, Respiro RSX01 dry powder inhaler). Attachable sensors are regarded as devices, whereas inhalers with embedded sensors are considered a drug with digital capabilities, and thus their approval process goes through different regulatory pathways. With each inhaler actuation, an electronic sensor time-stamps, records use, and stores data for a finite time. In addition to recording patient inhaler use, some manufacturers’ digital inhalers contain additional sensors that measure air flow during inhalation or report adequate shaking of pressurized metered-dose inhalers (pMDIs) prior to patient use (Table 1). These sensors rely on changes in pressure, sound waves, or vibration for such measurements. Built-in batteries typically last for the life of the inhaler, up to 13 months. In the United States, digital inhalers are required

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| Digital inhaler name | Description | Inhaler compatibility | Clinician interface | Patient interface | Pictures |
|----------------------|-------------|-----------------------|---------------------|------------------|---------|
| **Hailie® sensor range (formerly Smartinhaler™ sensors)** Adherium, Auckland, New Zealand | Records each actuation | Compatible with most pMDIs and DPIs (Diskus®, Turbuhaler®, HandiHaler®) | Portal dashboard reporting patient’s inhaler use | Smartphone dashboard | [Pictures](https://example.com) |
| **Respiro®** Amiko Milan, Italy | Records each actuation and inhalational flow for DPI and pMDI | Compatible with most pMDIs and DPIs (Diskus®, Turbuhaler®, HandiHaler®) | Web dashboard reporting patient’s inhaler use and inhalational flows | Smartphone dashboard | [Pictures](https://example.com) |

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Table 1 continued

| Digital inhaler name Manufacturer | Description Sensors | Inhaler compatibility | Clinician interface | Patient interface | Pictures |
|-----------------------------------|----------------------|-----------------------|---------------------|------------------|----------|
| INCA (Inhaler Compliance Assessment) | Records actuations and inhalational flows | Compatible with Diskus in UK | Portal dashboard reporting patient’s inhaler use and clinical data entered by patient | Smartphone dashboard Audible, scheduled alerts from sensor and/or smartphone | ![Picture 1](image1.png) |
| Dublin, Ireland | Attachable (bolt-on) sensor | | | | |
| Propeller sensors Propeller Health, Madison WI, USA | Attachable sensors Electromechanical | Compatible with most pMDIs, SMIs (Respimat®, Turbuhaler®, HandiHaler®, Ellipta®) Recently added Symbicort® pMDI | | Smartphone dashboard Environmental reports | ![Picture 2](image2.png) |
| | | | Alerts for worsening location of inhaler | | |
| Digital inhaler name | Description | Inhaler compatibility | Clinician interface | Patient interface | Pictures |
|----------------------|-------------|-----------------------|---------------------|------------------|---------|
| Digihaler            | Records each actuation and inhalational flow | USA                   | Portal dashboard reporting patient's inhaler use and inhalational flows (aid inhaler technique and physiological measure) | Smartphone dashboard | ![Image](image1.png) |
| Teva Pharm           | Embedded sensors measure pressure changes in airflow | ProAir Digihaler®      | Reports with trends of inhaler use and clinical data entered by patient | Audible, scheduled alerts from sensor and/or smartphone | ![Image](image2.png) |
| Tel Aviv, Israel     |                           | ArmonAir Digihaler (FP) |                                     | Alerts for worsening | ![Image](image3.png) |
|                      |                           | AirDuo Digihaler (FP/SAL) |                                     | Alerts for refills | ![Image](image4.png) |
|                      |                           |                       |                                     | Environmental reports | ![Image](image5.png) |
|                      |                           |                       |                                     | GPS based on location of inhaler | ![Image](image6.png) |
| HeroTracker sensor   | Bluetooth-enabled medication inhaler tracker sensors | pMDI and Diskus inhalers | Portal dashboard reporting patient's inhaler use | Smartphone and web dashboard: BreatheSmart® Connect—personal web dashboard for friends, family, and caregivers | ![Image](image7.png) |
| Cohero Health        |                           |                       | Reports for trends of inhaler use and clinical data entered by patient | BreatheSmart® app also available for reminder setting, inhaler tracking, lung function monitoring, and symptom recording | ![Image](image8.png) |
|                      |                           |                       |                                     | Links with mSpirometer lung function sensor (Bluetooth mobile spirometer) | ![Image](image9.png) |

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*DPI* dry powder inhaler, *FP* fluticasone propionate, *GPS* Global Positioning System, *PEF* peak expiratory flow, *pMDI* pressurized metered-dose inhaler, *SAL* salbutamol
to undergo manufacturer testing and meet regulatory standards depending on whether they are simply a sensor device (510(k) approval) or an inhaled drug with a built-in sensor [18].

Inspiratory-capable digital inhalers, which currently account for the minority of marketed products, can guide proper inhalation effort by patients to improve technique and could possibly serve as a physiological measure of lung function (Digihaler®, Respiro®). With the Digihaler®, measurement of inspiratory flow rate with each actuation allows the patient to see whether they achieved an adequate flow rate for drug delivery, but they currently are unable to view actual numbers for inspiratory flow. Clinicians can review the measured inhalational flow profiles to judge the patient's inhaler technique and to observe declines in lung function during exacerbations, and improvement in lung function as exacerbations are treated. The patient and clinician may see these inspiratory flow measurements as a value-added function when considering use of these devices.

A streamlined process allows a digital inhaler device to be registered and connected to a smartphone and cloud platform. Patients download the manufacturer-specific digital inhaler application onto their smartphone and then synchronize each inhaler through Bluetooth technology when both are in proximity. Each individual inhaler, whether the initial prescription or a refill, must be synchronized with the application. At the time of enrollment, platforms may ask patients to enter demographic and clinical data as well as details of the clinician and/or other caregivers accessing the data. After the data are entered, the patient and clinician can view their data on a dashboard accessed through their respective portals. The typical young person with asthma is likely to have a smartphone, whereas the older person with COPD or asthma may be less likely to possess one [19]. An alternative for the patient without a smartphone is to take their digital inhaler to an in-person clinic visit, and the data can be downloaded to a clinic computer using Bluetooth technology or by a USB connection.

Patients can interact with their smartphone digital inhaler application, including receiving daily controller reminder alerts, viewing current environmental conditions, recording their health status such as the Asthma Control Test (ACT) [20], and reviewing their inhaler use on a daily, weekly, or monthly basis. Devices provide audible and/or visual alerts to remind patients to use their maintenance inhalers daily; this function can be activated and de-activated by the patient. With some devices, the alert comes from the digital inhaler device, others through smartphone alerts. When there is rescue inhaler overuse, some platforms provide smartphone alert messages, advising patients to contact their clinician and health care team.

When the digital inhaler is synchronized with a smartphone that is connected to the internet, clinicians can access all data generated by these devices for each enrolled patient, with their consent. The clinician can, remotely and in real time, view the patient's controller adherence, assess some aspects of inhaler technique, identify over-users of medication, assess any clinical data provided by the patient on their digital dashboard, and generate reports of these events.

Key Roles of Digital Inhalers

The growing digital transformation of health care is revolutionizing the management of cardiac diseases, obstructive sleep apnea, and diabetes mellitus, among others [19]. Digital inhalers have been available for more than two decades but have mostly been employed in clinical studies, and uptake into routine clinical practice has been slow. More recently, with the expansion in the number of digital inhalers and their capabilities and the demonstrated benefits from their use in clinical trials, there is growing support among payers for their use in clinical practice [21]. Importantly, these devices are likely to lead to significant cost savings as a result of improved adherence and associated health outcomes in asthma and COPD [8, 22]. We discuss the important roles that these devices might play in the management of patients with asthma and COPD.
Characterizing and Improving Inhaler Adherence and Use

Medication adherence in patients with asthma and COPD is among the lowest for common chronic diseases [23], with significant impacts on health outcomes and costs [8, 22]. In both asthma and COPD, poor adherence has been associated with increased exacerbations, poorer symptom control, and mortality risk [6–8, 24, 25]. On average, long-term adherence with maintenance therapies averages 50% [26], though adherence in asthma and COPD may be even lower, as there are often no immediate consequences for the individual from nonadherence [27, 28]. Based on pharmacy claims data, regular use of daily maintenance medications is often < 50% in patients with asthma [29] and COPD [30]. This poor adherence and the resulting poor clinical outcomes has been shown to translate into increased direct and indirect costs, with studies reporting costs of US $7–17 billion being spent each year as a result of suboptimal inhaler use [31, 32]. Yet in patients with significant disease who are adherent to their inhaled medications, health care use is reduced, patient-reported outcomes (PRO) are improved, and mortality is decreased [27, 33, 34]. Studies in asthma and COPD support regular use of maintenance inhalers (termed adherent and controlled), and this is the basis for clinicians to promote daily use of controllers. Clinicians commonly overestimate adherence, and digital adherence monitoring will likely reveal that more of their patients are not well controlled and are not adherent to their prescribed regimen than clinicians currently recognize based on patient self-reports of inhaler use.

Digital Inhaler Use and Medication Adherence in Patients with Asthma

Digital inhaler use improves medication adherence, defined as regular daily use of controllers, and has resulted in reduced rescue inhaler use in several studies [35–43]. There are two types of monitoring by digital inhalers: passive, where biofeedback (BF) is provided to the patient and/or clinicians have access to the patient’s adherence data and are able to discuss them with the patient. Compared to passive monitoring of adherence, adherence is improved when BF is provided through reminder alerts and by active adherence monitoring by clinicians (Table 2) [35, 36, 39, 41, 43]. Active adherence monitoring would be defined as review of inhaler use either in real time or periodically such as with each clinical encounter. An RCT in 437 adults with uncontrolled asthma found that in the group provided BF using a smartphone application, adherence to fluticasone furoate/vilanterol dry powder inhaler (DPI) increased significantly, from a mean (standard deviation [SD]) percentage adherence of 82 (17)% over 24 weeks in the BF arm, compared to 71 (27)% in a control group with a passively monitored digital inhaler (study arm difference: 12%, 95% confidence interval [CI]: 5–19%; p < 0.001) [43]. When the clinician actively monitored adherence by reviewing patients’ inhaler use and providing feedback in real time, there was an additional 10% improvement. This modest increase should be interpreted with the caveat that adherence was much higher than normal in the control group. Other investigators reported greater improvements in controller adherence with digital inhalers [35, 37, 39, 44–46]. Another investigation in 100 adults with uncontrolled asthma found that over a period of 14 weeks, adherence to inhaled corticosteroids (ICS) declined less with the use of digital inhalers (−2%; 95% CI −7 to 3%; p = 0.40) than in the control group (−17%; 95% CI −26 to −8; p < 0.01) which did not receive reminders or feedback on medication use, representing a 15% (95% CI 4–25%; p < 0.01) difference [36]. At the same time, the percentage of short-acting beta agonist (SABA)-free days increased significantly in the digital inhaler group (19%; 95% CI 12–26; p < 0.01) versus a nonsignificant increase in the control group (6%, 95% CI −3 to 16; p = 0.18) [36].

In addition to validating pharmacy claims data, digital inhaler data provide greater granularity and accuracy, as claims data do not confirm actual medication use. A study in 1745 ambulatory children and adults with asthma
found adherence with controllers was optimal (> 80%) in 20%, moderately optimal (50 to ≤ 80%) in 28%, suboptimal (20–50%) in 25%, and poor (≤ 20%) in 27% [53]. These investigators reported that adherence with controllers declined by more than 25% over 20 weeks, even in patients initially optimally adherent [53]. The largest decrease in adherence was observed for the moderately adherent group (~32%) and least for the poor adherence group (~6%), likely because of the already low level of adherence. The poor adherence group also had an 8.1% (95% CI 2.9–13.3%; p < 0.01) higher proportion of patients prescribed ICS only compared to those in the other adherence groups (63.7% vs. 55.6%). This finding suggests that an ICS product containing a bronchodilator may support better adherence compared to ICS therapy alone. In a feasibility study of children with asthma after discharge from the emergency department (ED), who relied on reminder alerts but did not employ active digital monitoring, adherence to controllers was reported to be only 36% in an intervention group compared to 32% in the control group [54].

**Digital Inhaler Use and Medication Adherence in Patients with COPD**

Similar to patients with asthma, regular use of maintenance inhalers occurs in only about half of patients with COPD [41, 55]. In a high-risk population of post-hospitalized patients with COPD, where daily use of controllers is certainly justified, adherence (when erroneously taken events were excluded) was only 30% over 1 month when measured using a digital ICS/long-acting beta agonist (LABA) DPI provided at discharge. There was wide variation in actual use, as < 20% of patients used their inhaler regularly and with the correct technique [41].

**Digital Inhalers Identify Patterns of Medication Use**

Patterns of rescue inhaler use have been better characterized with digital technology [56–58]. In a subpopulation of the COPDGene study (n = 58), four SABA patterns were identified using a digital inhaler: (1) frequent use, regular pattern, (2) frequent use, no pattern, (3) infrequent use, (4) infrequent, but intense use [57]. Groups 2 and 3 were the most common patterns. As expected, albuterol use was driven by respiratory symptoms. In a prospective, observational study of 32 patients with COPD who overused albuterol, 73% were on maximal therapies, while only 27% of those with overuse were on suboptimal therapies [58]. This study also found that albuterol use was strongly associated with symptoms [58]. Neither study looked at adherence to controllers [57, 58]. Using digital inhalers to obtain data on patterns of medication use is also key to identifying an individual’s reasons for SABA overuse or poor controller adherence, for example; the clinician can review inhaler use data with the patient to identify any recurring triggers for increased SABA use or ICS underuse, such as job commitments or travel. This represents an important opportunity to improve adherence by addressing modifiable sources of nonadherence; similarly, this allows clinicians to also identify individuals who are intentionally non-adherent, for example, due to concerns about side effects, for whom a more intensive, health psychology-based intervention may be needed.

The importance of understanding patterns of SABA use is reflected in the current asthma guidelines [59, 60] and PROs such as the ACT [61] and SABA Reliance Questionnaire (SRQ) [62]. These include measures of SABA frequency to judge disease control and/or severity, and risk of over-reliance on SABA. The SRQ assesses patient perceptions of SABA and identifies key beliefs that may be driving SABA overuse [62]. The frequency of β-2 agonist use is commonly used by clinicians during encounters, and assessment of SABA use is now recommended in current guidelines as a measure of disease control for asthma or COPD, as overuse of SABA is related to increased morbidity and mortality [56]. Documenting rescue SABA use with digital inhalers could help clinicians more accurately determine disease severity for asthma and predict asthma exacerbations [56, 63].

Digital inhalers can also be used to evaluate patterns of controller medication use, similarly to informing about SABA use patterns [64]. Previous studies using digital inhalers have identified different patterns of adherence,
| Author (year)          | Study population                                                                 | Digital device | Patient and clinician interfaces with EMD and apps | Primary endpoint(s)                |
|-----------------------|----------------------------------------------------------------------------------|----------------|---------------------------------------------------|------------------------------------|
| Alshabani (2018) [47] | COPD patients at high risk of adverse events from COPD (n = 20) 6 months       | Propeller (albuterol pMDI) | EMD with clinician dashboard                      | All-cause HCU with EMD + dashboard compared to prior year |
| (Abstract only)       |                                                                                  |                 | Poor compliance by excess SABA or daily controller use prompted clinician response |                                    |
| Alshabani (2020) [48] | COPD patients with high health care utilization (n = 39) 12 months               | Propeller (digital inhalers for both controller and albuterol pMDI) | EMD with patient dashboard                      | All-cause and COPD HCU compared with year prior to enrolment in study |
| Retrospective pre- and post-analysis, open-label |                                                                                   |                 | Patients alerted when adherence to controller was suboptimal or rescue inhaler use increased |                                    |
| Barrett (2017) [40]   | People with asthma > 4 years old (n = 95) 13 months                               | Propeller       | Smartphone app to aid patient self-management     | SABA use compared to 30 day baseline |
| Prospective, observational Community project |                                                                                   | Albuterol pMDI | No clinician dashboard                           | ACT and c-ACT compared to 30 day baseline |
| Chan (2015) [35]      | Children with asthma aged 6–15 years on ICS with recent exacerbation requiring ED (n = 220) | Adherium        | Patient adherence reminders by device             | School absenteeism (co-primary)     |
| RCT                   |                                                                                  | SmartTrack® ICSPMDI | Medication adherence (co-primary)                 |                                    |
|                       |                                                                                  |                 |                                                   |                                    |
| Foster (2014) [37]    | Adults with mod/severe asthma based on ACT (n = 143)                              | Adherium        | Patient inhaler reminders with clinician dashboard prompting patient contact with suboptimal inhaler adherence | ACT                               |
| RCT                   |                                                                                  | SmartTrack® ICS/LABA (Accuhaler®) |                                             | Mini-AQLQ                          |
| 6 months              |                                                                                  |                 |                                                   | Anxiety/depression scale           |
|                       | EMD + BF + clinician/patient adherence discussions vs. EMD + BF vs. Clinician/patient adherence discussion only vs. Usual care 6 months | Albuterol pMDI |                                                   |                                    |
| Author (year) | Study population | Digital device | Patient and clinician interfaces with EMD and apps | Primary endpoint(s) |
|---------------|------------------|----------------|-----------------------------------------------|---------------------|
| Gregoriano (2019) [49] | Adult asthma and COPD with exacerbation in last year (n = 149) | Adherium Hailie® sensor (formerly SmartInhaler) | EMD with clinician dashboard + BF (patient inhaler alerts, clinician assessments) | Time to first exacerbation |
| Single-blind RCT, Pulmonary clinic | 6 months | Albuterol pMDI | vs. Passive EMD | |
| Kaye (2020) [42] (Abstract only) | COPD (n = 1000) | Propeller | EMD with patient dashboard | CAT at 6 months compared to baseline |
| Prospective, observational | 6 months | Albuterol pMDI | | Albuterol use |
| Lin (2020) [50] | Asthma 10–17 years old (n = 21) | Propeller Health | Monthly clinic visits | Composite Asthma Severity Index (CASI) |
| Open-label study, Inner city school-based intervention, Inner city school | 6 months | Albuterol pMDI and controllers | Clinician and patient dashboards | |
| Merchant (2016) [38] | Children and adults with asthma > 5 years old (n = 495) | Propeller | Clinician access to dashboard | SABA-free days |
| RCT, parallel arms, Allergy clinic | EMD + BF vs. Usual care, 12 months | Albuterol pMDI | Personalized feedback to patient via mobile phone app | |
| Merchant (2018) [51] | Adults with asthma (n = 224) (n = 76 on controller medication) | Propeller | Clinician access to web-based dashboard | ED visits |
| Prospective, open-label, Allergy clinic | 12 months | Albuterol pMDI | Patient feedback from software on personal device | Hospitalizations |
| | | Various controllers | Combined ED visits and hospitalizations | |
| Author                  | Design           | Setting                | Study population                              | Digital device Drug(s) | Patient and clinician interfaces with EMD and apps |
|-------------------------|------------------|------------------------|-----------------------------------------------|------------------------|---------------------------------------------------|
| Moore (2020) [43]       | (Abstract only)  | Open-label, parallel-group RCT | Adults with uncontrolled asthma (ACT < 20) on fixed-dose ICS/LABA maintenance therapy (n = 437) | Ellipta sensor Fluticasone furoate/vilanterol Ellipta DPI Albuterol pMDI | One of five connected inhaler systems with different levels of data feedback from sensors: (1) Maintenance use to participants and health care professionals (HCPs) (N = 87); (2) Maintenance use to participants (N = 88); (3) Maintenance and rescue use to participants and HCPs (N = 88); (4) Maintenance and rescue use to participants (N = 88); (5) No feedback (control) (N = 86) |
| Morton (2017) [46]      | Open-label RCT   | Pediatric asthma 6–16 years old (n = 77) | Intervention group (EMD with controller reminders and review of adherence at clinic visits) | Adherium Hailie® sensors for pMDI and for turbuhaler (formerly Smartinhale®, SmartTurbo®) ICS pMDI and ICS DPI SABA pMDI | Passive recording of EMD use Adherence reviewed at clinic visits with health care provider Asthma Control Questionnaire |
| Author (year) | Design | Setting | Study population | Digital device | Patient and clinician interfaces with EMD and apps | Primary endpoint(s) |
|---------------|--------|---------|------------------|----------------|----------------------------------------------------|---------------------|
| Mosnaim (2020) [36] | RCT, single blinded | Allergy clinic | Adults with uncontrolled asthma \((n = 100)\) vs. Usual care with passive EMD | Propeller | Yes | SABA-free days |
| O'Dwyer (2020) [39, 52] | Randomized parallel study | Community pharmacies and clinics | Adult asthma \((n = 83)\) vs. COPD \((n = 55)\) | INCA (Seretide® Accuhaler) | EMD + BF | ICS/LABA adherence (frequency of use and proper inhaler technique) |
| Sulaiman (2018) [41] | RCT | Pulmonary clinic | Adults with severe asthma with exacerbation in last year \((n = 360)\) vs. Control (Intensive education) | INCA (Seretide®) | | ICS/LABA adherence (frequency of use and inhaler technique) |
| Author (year) | Secondary endpoint(s) | Primary outcome | Secondary outcome | Comments |
|--------------|------------------------|----------------|-----------------|----------|
| Alshabani (2018) [47] | (Abstract only) | Prospective open-label | All-cause HCU (hosp + ED) lower with EMD + BF ($p = 0.034$) | COPD-related hospitalization was also lower with EMD + BF (2.3 (2.2) vs. 3.9 (3.2) but not significant ($p = 0.07$), no CI provided) | Prompted patient contact by clinicians |
| Alshabani (2020) [48] | Reducing in COPD-related HCU per year (2.2 ±2.3 vs. 3.4 ±3.2, $p = 0.01$). Reduction in all-cause HCU, but this was not statistically significant (3.4 ±2.6 vs. 4.7 ±4.1, $p = 0.06$), no CI provided | Average adherence was 44.4% (28.4%), with mean Charlson comorbidity index 5.6 (2.7) |
| Barrett (2017) [40] | --- | Prospective, observational | 39% ↓ in SABA use in first 30 days ($p < 0.001$) | Increase in percentage of symptom-free days from 77% during the baseline period to 86% after the first month (12% improvement) | Higher proportion with controlled asthma |
| Author (year) | Design | Setting | Study population | Study group(s) | Duration | Digital deviceDrug(s) | Patient and clinician interfaces with EMD and apps | Primary endpoint(s) |
|--------------|--------|---------|------------------|----------------|----------|----------------------|-------------------------------------------------|-------------------|
| Chan (2015) [35] | RCT | Children with asthma presenting to ED | Asthma morbidity score | Child ACT | Asthma morbidity score | Median medication adherence: 84% (10th percentile 54%, 90th percentile 96%) with EMD vs. 30% (8%, 68%) in control group \( p < 0.0001 \) | Improved asthma score from baseline with EMD | Adherence reduced over 6 months in both groups though difference between two groups remained statistically significant |
| | | | Child ACT | FEV1 | Exacerbations | Improvement in asthma morbidity score \( p = 0.008 \) and Childhood ACT \( p < 0.0001 \) improved in EMD + BF | Lower SABA use in EMD + BF (median % of days of SABA use = 9.9% [10th percentile 1.1%, 90th percentile 32.8%] in the intervention group vs. 17.4% [2.4%, 49.2%] in the control group; \( p = 0.002 \)) | No effect on FEV1 |
| Foster (2014) [37] | RCT | 6 months | ICS/LABA adherence | FEV1 | No difference in ACT among 4 groups \( p = 0.14 \) | No difference in other PRO or FEV1 among 3 groups |
| | | | | | | | |

360 Pulm Ther (2021) 7:345–376
| Author (year) | Design | Setting | Study population | Study group(s) | Duration | Digital device | Drug(s) | Patient and clinician interfaces with EMD and apps | Primary endpoint(s) |
|--------------|--------|---------|------------------|---------------|----------|---------------|---------|-------------------------------------------------|-------------------|
| Gregoriano (2019) [49] | Single-blind RCT | Pulmonary clinic | Exacerbation frequency | Controller adherence | SGRQ score | No effect on time to first exacerbation (HR 0.65, 95% CI: 0.21–2.07, \( p = 0.024 \)) | Nonsignificant decrease in exacerbation frequency (RR = 0.61, CI: 0.35–1.03, \( p = 0.07 \)) | Days adherent greater in intervention group |
| Kaye (2020) [42] | Prospective, observational | | Exacerbations defined by acute worsening requiring contacting the clinician (not based on oral steroid use) | | | Albuterol use declined by 0.8 puffs/day (95% CI: \(-0.9\) to \(-0.7\); \( p < 0.001 \)) compared to baseline | Albuterol use greater in non-adherent patients | 46% of subjects met MCID for CAT (\( \geq 2 \) change) |
| Lin (2020) [50] | Open-label study | Inner city school | School absenteeism | Exacerbations | Medication adherence | No change in mean CASI from baseline; no CI provided | Decrease in school absenteeism (\( p = 0.003 \)) | 81% with step-up in controller therapies |

Exacerbations defined by acute worsening requiring contacting the clinician (not based on oral steroid use)
| Author (year) | Design | Setting | Study population | Study group(s) | Duration | Digital device | Drug(s) | Patient and clinician interfaces with EMD and apps | Primary endpoint(s) |
|--------------|--------|---------|------------------|----------------|----------|----------------|---------|---------------------------------------------------|-------------------|
| Merchant (2016) [38] | RCT, parallel arms | Allergy clinic | ACT | EMD + BF vs. usual care group | 17% vs. 21%, p < 0.01; no CI provided | † in ACT with EMD + BF vs. control group (+6.2 vs. +4.6, p < 0.01) | Attrition 14.8% at 1 month and > 55% at 12 months |
| Merchant (2018) [51] | Prospective, open-label | Allergy clinic | SABA use | ICS adherence | ED visits lower compared to baseline period (rate difference 6.3 [95% CI: 0.9–11.6], p = 0.04) | SABA use ↓ by 0.52 puffs/daily (95% CI: −0.69 to −0.34; p < 0.05) | ↓ inpatient visits by 2.6 (95% CI: 2.2, 2.9) visits per patient-year (p < 0.01) |
| Moore (2020) [43] (Abstract only) | Open-label, parallel-group RCT | SABA-free days | ACT score | Mean (SD) adherence 82.2 (16.58)% in the "maintenance to participants and HCPs" arm and 70.8 (27.30)% in the control arm; difference of 12.0% (95% CI: 5.2–18.8%, p < 0.001) | Mean % SABA-free days (months 4–6) significantly greater in those who received data on rescue use vs. control arm | Only measured inhaler use months 4–6 |
| Author (year) | Design | Setting | Study population | Study group(s) | Duration | Digital device | Drug(s) | Patient and clinician interfaces with EMD and apps | Primary endpoint(s) |
|--------------|--------|---------|-----------------|----------------|----------|---------------|---------|------------------------------------------------|-------------------|
| Morton (2017) [46] | Open-label RCT | Acute care visits to office or ED | Adherence to ICS and use of SABA | FEVI | Pediatric Asthma QOL Questionnaire (PAQLQ) | Guideline-based severity | No difference in Asthma Control Questionnaire (difference of the paired mean difference from baseline to 12 months = −0.18 (95% CI: −0.76 to 0.38; p = 0.51) | Adherence improved in intervention group 70% vs. 49% (p = 0.001); no CI provided | Fewer exacerbations in intervention group (controls 53% more likely to receive steroids—incident risk ratio = 1.53 [95% CI: 1.11–2.11]; p = 0.008) and hospitalizations (p < 0.001) No significant difference in SABA use, FEVI, or PAQLQ or asthma severity |
| Mosnaim (2020) [36] | RCT, single blinded | Allergy clinic | ICS adherence | | | | | | | Fewer exacerbations and asthma control not different |

Exploratory outcomes of exacerbations and asthma control not different
| Author (year) | Setting | Study population | Study group(s) | Duration | Digital device | Drug(s) Patient and clinician interfaces with EMD and apps | Primary endpoint(s) |
|--------------|---------|------------------|----------------|----------|----------------|----------------------------------------------------------|-------------------|
| O’Dwyer (2020) [39, 52] | Community pharmacies and clinics | Randomized parallel study | Resp symptoms | BF in inhaler instruction vs. vs. | EMD + BF + inhaler instruction vs. Inhaler training group had improvement at 2 months, but not at 6 months | SGRQ (−6.1; 95% CI: −9 to −0.4; p = 0.04) in EMD + BF group at 2 and 6 months | Pharmacies were unit of randomization |
| Sulaiman (2018)[41] | Pulmonary clinic | ACT | 73% adherence (frequency of use, correct technique) in EMD + BF vs. 63% in control group (difference = 10%; 95% CI: 2.8–17.6%; p = 0.02) | PEF at 3 months, no statistically significant difference between groups | ACT or AQLQ, no statistically significantly different between groups | 27% of participants with severe asthma were refractory to current treatment despite being adherent and receiving intensive education |

**Table 2 continued**

**Notes:**
- ACT Asthma Control Test
- AQLQ Asthma Quality of Life Questionnaire
- BF = biofeedback
- CAT COPD Assessment Test
- CI confidence interval
- COPD chronic obstructive pulmonary disease
- DPI dry powder inhaler
- ED emergency department
- EMD electronic monitoring device
- FEV1 forced expiratory volume in 1 second
- HCU health care utilization
- ICS inhaled corticosteroids
- LABA long-acting beta agonist
- MCID minimal clinically important difference
- PAQLQ Pediatric Asthma Quality of Life Questionnaire
- PEF peak expiratory flow
- pMDI pressurized metered-dose inhaler
- PRO patient-reported outcomes
- RCT randomized controlled trial
- SD standard deviation
- SABA short-acting beta agonist
- SGRQ St George’s Respiratory Questionnaire
| Author (year) | Study population | Digital device | Outcome measures | Outcome | Comments |
|--------------|------------------|---------------|------------------|--------|----------|
| Killane (2016) [69] | Adults with asthma ($n = 184$) | INCA | Exacerbation risk | < 80% adherence by EMD predictive of adverse events of COPD | Digital records of adherence more accurate than relying on dose counter. Both predictive of exacerbations |
| Pleasants (2019) [74, 75] | Adults with asthma ($n = 360$) on ICS/LABA with exacerbation in prior year | Teva Digihaler® Albuterol | Clinical, β-agonist use, and inspiratory flow measures to predict exacerbations using machine learning modeling | PIF and inhalation volume measured by Digihaler decline with exacerbations Albuterol use increases with exacerbations | On average, patients without exacerbations used ProAir Digihaler 1.17 (SD = 1.51) times per day vs. 1.82 (2.13) for those who had 1 or more exacerbations (outside the exacerbation period) |
| Snyder (2020) [76, 77] | COPD with history of exacerbation ($n = 336$) | Teva Digihaler® Albuterol DPI | Clinical, β-agonist use, and inspiratory flow measures to predict exacerbations | PIF and inhalation volume measured by Digihaler decline with exacerbations Albuterol use increases with exacerbations | |
| Sumino (2018) [78] | COPD ($n = 35$) | Propeller Albuterol pMDI Passive EMD without dashboard | Exacerbation risk based on albuterol use compared to baseline | Odds ratio of an exacerbation 1.54 (95% CI: 1.21–1.97 with ↑ albuterol use > 100%) | |
ranging from regular adherence and irregular adherence, to regular nonadherence and irregular nonadherence [64, 65]. Clinicians will likely encounter patients who are non-adherent but controlled, though little has been published in this area.

Digital Inhaler Use in Clinical Trials

Digital inhalers could also be employed in clinical drug trials to quantify inhaler adherence and lessen the impact of nonadherence on study results [66, 67]. Similarly to the clinical setting, drug trials are heavily dependent on patient self-reporting before and during the

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Table 3 continued

| Author (year) | Study population | Digital device | Outcome measures | Outcome | Comments |
|---------------|------------------|----------------|------------------|---------|----------|
| Patel (2013) [56] | Severe asthma \(n = 303\) | Adherium | Effect of albuterol use to predict exacerbations (used first 2 weeks of study period to define baseline use) | Each associated with an increased risk of future severe exacerbation | Higher mean daily albuterol use \(OR 1.24; 95\% CI: 1.06–1.46\) |
| Hoch (2019) [63] | Adults with asthma or COPD \(n = 2509\) asthma and 899 COPD | Propeller | Linear model to predict high use of albuterol (peak SABA use \(\geq 6\) puffs/day) | ↑ Albuterol use by 100% for 3 days predictive of the event \(p < 0.01\) | Higher SABA use \(p < 0.01\) and higher variation in use (2.5–97.5th percentiles) was observed in the post-vs. pre-SABA use peak period, and 41–43% had \(\geq 1\) additional high-use day within 10 days after the initial peak event |

AE adverse events, CI confidence interval, COPD chronic obstructive pulmonary disease, DPI dry powder inhaler, EMD electronic monitoring device, FOR formoterol, ICS inhaled corticosteroids, LABA long-acting beta agonist, OR odds ratio, PIF peak inspiratory flow, pMDI pressurized metered-dose inhaler, RCT randomized controlled trial, SABA short-acting beta agonist
study, with many using daily diaries to document medication use. However, self-report measures are prone to overestimation of adherence due to social desirability bias [68]. Dose-counters on inhalers are also used to measure adherence in trials but have been found to overestimate actual use compared with the use of a digital inhaler in practice [69]. While run-in periods and study design can lessen the influence of inhaler nonadherence on outcomes, the actual impact is unknown without the use of these digital devices throughout the study. There is great potential in using these devices to better understand the “true” relationship between medication dose and efficacy, for example in quantifying how different patterns of use (different doses and dosing frequency) relate to reported outcomes [70]. Whilst there have been studies funded by pharmaceutical companies which explore the use of digital inhalers and impact on adherence and outcomes, such as the effect of an add-on device to a Turbuhaler [71] or smartphone application plus device added to a pMDI [72], these have been limited to evaluation of the digital device itself as an adherence intervention, rather than using digital inhalers to explore drug dose efficacy relationships as part of the clinical drug trial. A recent systematic review of published clinical trials of asthma add-on step 4/5 therapy assessed how prior adherence to inhalers had been assessed. In this review, it was identified that none of the 87 studies had used objective measures to assess adherence, before or during the trials. In some modeling work, the authors estimated that this lack of assessment increased the statistical variance of the outcomes and described how this led to much larger than needed sample sizes [73].

Role in Reducing Frequency of Exacerbations

Digital inhalers collect rescue and controller data virtually and in real time, track adherence, and often include clinical platforms that aid patient self-management; their use would be expected to better prevent exacerbations. Five studies in asthma or COPD found lower exacerbation rates, two as the primary outcome [48, 51] and three as a secondary outcome (Table 3) [37, 46, 49]. These studies used some form of active adherence monitoring by the clinicians. Two other studies in asthma or COPD, one with passive [35] and the other with active monitoring [36], found no difference in exacerbation rates. In the study by Chan and colleagues, frequency of exacerbations decreased during the initial 2 months, but this difference was lost at 4 and 6 months as adherence trended towards lower levels [35]. A 6-month RCT in 149 asthma and COPD patients did not show a significant difference in time to first exacerbation using digital monitoring of rescue and controller medications with biofeedback (BF) (hazard ratio [HR] 0.65, 95% CI 0.21–2.07, \( p = 0.24 \)), but there were fewer exacerbations in the EMD + BF group than in the control group, although this was not statistically significant (36.6% vs. 63.3%, relative risk [RR] = 0.61, 95% CI 0.35–1.03, \( p = 0.07 \)) [49]. The use of digital rescue inhalers was studied in 224 adult patients with asthma followed for 1 year. Patients received digital inhalers that tracked rescue and controller inhaler medication use, and a digital health platform that presented medication use information and asthma control status to patients and providers. Decrease in hospitalizations did not meet statistical significance (exacerbation rate 1.8 [95% CI 0.5–4.6] pre-enrollment vs. 0.4 post-enrollment [95% CI 0.01–2.5]; rate difference 1.3 [95% CI −0.6 to 3.3], \( p = 0.23 \)), but ED visits alone (11.6 vs. 5.4; rate difference 6.3 [95% CI 0.9–11.6], \( p = 0.04 \)) and combined ED and hospitalizations (13.4 vs. 5.8; rate difference 7.6 [95% CI 1.9–13.3 \( p = 0.02 \)) were both significantly lower with the use of digital rescue inhalers compared with baseline levels before enrollment in the study [51].

In addition to reducing exacerbation risk by improving controller adherence, digital inhalers can identify increases in rescue inhaler use that are associated with exacerbations. In adults with asthma, increased rescue inhaler use with budesonide/formoterol was reported about 5 days prior to an exacerbation, and then use declined over a similar interval as the
exacerbation was treated [79]. Previously, this pattern was also described using self-report of rescue albuterol use in patients with asthma [80]. Two groups of investigators found that an increase in SABA use by 100%, as recorded by digital inhalers, was predictive of an ensuing exacerbation [63, 78].

Inspiratory-capable digital inhalers add a physiological measure to the drug usage data around exacerbations. Similar to peak expiratory flow, peak inspiratory flow (PIF) is responsive to bronchodilators [81] and decreases during exacerbations [74, 76, 82]. Air trapping and airflow obstruction are important mechanisms for reducing PIF [83, 84]. The Digihaler® and Respiro® are digital inhalers that can measure PIF, inhalation volume, and time to peak. Three studies have reported acute changes in PIF around exacerbations, two using digital DPIs. In hospitalized patients with COPD or asthma, Chrystyn and colleagues found that PIF measured with a portable inspiratory flow meter was low upon admission (often < 30 L/min), and increased over several days [83]. In a study of 360 adults with asthma using the Digihaler®, 64 of whom experienced an exacerbation, PIF began decreasing about 5 days prior to the exacerbation, then returned to baseline over a similar interval [74]. A study of the ProAir Digihaler® in 336 patients with COPD, of whom about one-third (n = 98) reported an exacerbation, found that PIF had not returned to baseline by 14 days, similar to expiratory flow patterns reported post-exacerbation [85]. By using machine learning for both trials, increased SABA use was the strongest predictor of an impending exacerbation in asthma, while in COPD, inspiratory flows were most important [76, 77]. This has important implications when considering the role of digital inhalers, as inhalers that measure PIF could play a key role in asthma management by predicting risk of impending exacerbation through a reduction in PIF and inhalation volume or by recording more frequent SABA use. Additionally, monitoring PIF may help patients identify when inspiratory flow rates are too low for adequate drug delivery from DPIs, informing them to switch to nebulized or pMDI therapy, as the minimal inspiratory flow rate required for effective treatment from DPIs is at least 30 L/min [86].

Improving Inhaler Technique

All digital inhaler devices can identify instances where patients administer their doses too closely together. This “dose dumping” phenomenon may reflect an important source of technical errors in the use of pMDIs, or in clinical trials may reflect social desirability bias from participants wishing to “window-dress” their adherence, or to conceal nonadherence, prior to a study visit or observation [35, 87]. A study using Propeller Health devices found that 67% of patients administered doses of SABA pMDI within 15 seconds of each other, suggesting that many patients inhale too rapidly and/or do not have adequate breath-hold [88], or that they are deliberately attempting to conceal nonadherence [35]. Inspiratory-capable digital DPIs (Digihaler®, INCA®, Respiro®) can guide proper inhaler technique by recording whether minimum desired flows are achieved. For DPIs (Digihaler®, INCA®, and Respiro®), thresholds for PIF are used to determine whether the minimal PIF is achieved, but there is inconsistency in the literature as to what is a minimally acceptable PIF for individual DPIs [89]. In contrast to DPIs, a slower inspiratory flow rate is necessary for optimal drug delivery from pMDIs. The Respiro® add-on device for pMDIs is one such digital device that measures whether the inhalation is slow enough. Other digital devices are expected to provide similar assessments of pMDIs [14].

Published data concerning proper inhaler technique primarily involve the INCA device (fluticasone propionate [FP]/salmeterol [SAL] DPI [Accuhaler®]) in patients with asthma or COPD [41, 90]. Among post-hospitalized patients with COPD, multiple errors in inhaler technique occurred during a 1-month observation period. These included opening and closing the inhaler without using the device, blowing into the device, and most commonly poor inspiratory effort [90, 91]. Only 24% of individuals used their inhaler with the proper technique and at the correct time interval as
prescribed. The same investigators also reported on inhaler technique using INCA in adults with severe asthma over 3 months [41]. This study compared digital FP/SAL use with intensive patient education and BF from the clinician with a comparison group who received digital FP/SAL and intensive education without BF. There was no significant difference in the proportion of patients with proper inhaler technique; however, subjects did not receive immediate feedback on their technique or inhaler use. Thus, we are unaware of published evidence that patient use of digital inhalers improves inhaler technique when compared to intensive teaching alone or even to usual care.

**Enhancing Patient-Reported Outcomes**

Seven studies have reported on the impact of digital inhalers on PROs in childhood asthma [35, 38, 40, 46], adult asthma [36–38, 40, 41, 49, 52], and COPD [42, 49, 52] (Table 2). Overall, PROs improved in most studies in patients with asthma whether with passive [35, 40, 41] or with active clinician monitoring [36, 38, 42, 52]. In two studies in patients with asthma, improvements in PROs were not significant [37, 46]. Interestingly, PROs improved when digital inhalers with SABA alone were employed (i.e. there was no controller treatment) [38, 40, 42], perhaps owing to closer monitoring and collaboration by the research/health care team. PROs improved with the use of digital inhalers in two studies in patients with COPD [42, 52], but not in another study [49], possibly because the latter study included only those patients who had suffered an exacerbation in the previous year, thus possibly representing patients who might have had more uncontrolled disease compared to the other two studies.

**Improvement in Pulmonary Function**

A few investigators have reported an improvement in lung function with the use of digital inhalers in people with asthma [35, 37, 41, 46], but only as a secondary study endpoint. The impact of digital inhaler use on pulmonary function in patients with COPD has not been reported. In two studies among pediatric asthma patients [35, 46], no significant changes were found in FEV1 using either digital SABA or controller inhalers; neither employed active monitoring by clinicians. In a study in adults with asthma that used active monitoring by clinicians, no significant effect on FEV1 [37] or peak expiratory flow (PEF) was found [41], despite a significant improvement in adherence to controller medications [35, 37, 41, 46]. However, pulmonary function may not accurately reflect asthma control, and other measures reflective of asthma control may need to be evaluated [92].

**Informing Costly and Potentially Risky Interventions**

Various therapeutic strategies are increasingly being used for asthma and COPD patients who fail to achieve adequate control with inhaled medications. These include bronchial thermoplasty, biologics for asthma, and airway stents for COPD. In a study of adults with uncontrolled asthma who were considered eligible for biologics or bronchial thermoplasty, more than half were non-adherent with their digital controller inhalers [93]. Another interesting observation from this study was that only 5% of subjects referred to the specialty clinic for uncontrolled asthma were thought to be non-adherent by the referring physician. The uncertainties around COPD and asthma diagnosis further complicate this, particularly when data suggest that up to 30% of patients on inhalers may not have an asthma diagnosis when objective diagnostic measures are applied [94]. Ensuring correct diagnosis and evaluating adherence using digital inhalers are important steps to conduct prior to adding very expensive biologics to patients’ treatment regimens. Using digital inhalers to gather more information on patient medication-taking patterns may be the most impactful cost-effective use of these devices. A recent economic analysis of digital inhaler use in COPD patients suggested that using digital inhalers to tailor and target interventions based on personal adherence patterns may
be cost-effective and potentially cost-saving [95]. For example, digital inhalers can be used to assess environmental triggers of asthma, where data on reliever use aligned with geolocation can inform identification of triggers [96] to alert the patient to potential exposures, thus potentially preventing worsening asthma and/or costly exacerbations. Targeting the use of digital inhalers to those who are most likely to benefit from treatment will increase their cost-effectiveness [35].

**Considerations for Implementation and Future Applications**

Cost, patient and provider acceptability, data management, sustainability of digital technology, and effective integration into the care pathway remain key barriers to uptake of this technology into routine clinical practice [97]. Delineating the role of digital inhalers, for example, in diagnosis and disease monitoring can help improve uptake. Current research is underway to identify patient characteristics of those who may most likely benefit from the use of digital inhaler-based interventions [98]. Additionally, there are ethical issues for patients, providers, and other stakeholders that may need to be considered prior to implementation into practice, such as informed consent, autonomy, trust, privacy, confidentiality, and remote patient monitoring [99]. These issues, however, are not specific to digital inhalers; in other areas of medicine, such as cardiovascular disease, diabetes, and tuberculosis, remote patient monitoring has been around for a long time, and these ethical and privacy issues have been well addressed and discussed [100], including consideration of the unique benefits that electronic adherence monitoring brings. Specifically, in terms of data collection, digital inhalers only collect data specific to the device, primarily the time and date of each dose, with some devices logging PIF (e.g. Digihaler). Location data are only collected when paired with an app; the devices themselves do not log or record other data. Even audio-based devices record data for less than a minute and then “time out” after that, and the microphone is placed to avoid ambient sounds or other unrelated audio. In terms of cloud management of data, similar to other electronic medical records, individuals can choose to opt out of cloud sharing of data and only share data via direct download between the device and computer; any data that are transferred on the cloud should be Health Insurance Portability and Accountability Act (HIPAA)-compliant [101]. Digital inhalers have the advantage of providing detailed information about medication taking which, when combined with artificial intelligence approaches, can revolutionize asthma and COPD care. For example, InspirerMundi allows remote monitoring of inhaler use via a machine learning approach that can assess medication self-administration [102]. If these artificial intelligence approaches can be applied to the “big data” captured by digital inhalers, there is potential that the data can help predict and/or mitigate future exacerbations. However the amount of data required to do this will need to be large and likely need to be informed by multiple data sources, beyond that captured by the inhaler device for example, linking with data collected from smartphones, digital spirometers, and/or other wearables such smartwatches [101].

**CONCLUSION**

Despite their availability for over two decades, digital inhalers and associated health management platforms remain new and unfamiliar to many clinicians and health care organizations. Typically, patients use their smartphones for data transfer and health management, providing remote access to the clinician. Available evidence indicates that digital inhalers enhance medication management and guide clinical care in patients with asthma or COPD, with benefits of increased medication adherence having the potential to improve clinical outcomes and prevent the need for costly or more risky therapies, such as biologics. However, evidence regarding acceptability for patients and end-users, cost-effectiveness, improvement in inhaler technique, and the best practice models to integrate these devices into routine care

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remains somewhat elusive. While studies support several benefits of digital inhalers, there are many unanswered questions, including the extent to which they will be adopted in clinical practice.

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