The COVID-19 Pandemic and Eating Disorders in Children, Adolescents, and Emerging Adults: Recommendations from the Canadian Consensus Panel

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ABSTRACT

Objective: The COVID-19 pandemic has had detrimental effects on mental health. Literature on the impact on individuals with eating disorders is slowly emerging. While outpatient eating disorder services in Canada have attempted to transition to virtual care, guidelines related to optimal virtual care in this field are lacking. As such, the objective of our Canadian Consensus Panel was to develop clinical practice guidelines related to the provision of virtual care for children, adolescents, and emerging adults living with an eating disorder, as well as their caregivers.

Methods: Using scoping review methodology (with literature in databases from 2000-2020 and grey literature from 2010-2020), the Grading of Recommendations, Assessment, Development, and Evaluation system, the Appraisal of Guidelines, Research and Evaluation tool, and a panel of diverse stakeholders from across Canada, we developed high quality treatment guidelines that are focused on virtual interventions for children, adolescents, and emerging adults with eating disorders, and their caregivers.

Results: Strong recommendations were supported specifically in favour of in-person medical evaluation when necessary for children, adolescents, and emerging adults. For children and adolescents, weak recommendations were supported for telehealth family-based treatment (FBT) and online guided parental self-help FBT. For emerging adults, cognitive-behavioural therapy (CBT)-based virtual group therapy and internet CBT-based guided self-help, as well as internet-based Maudsley Model of Anorexia Nervosa Treatment for Adults (MANTRA) guided self-help were strongly recommended. Weak recommendations for emerging adults included telehealth relapse prevention using MANTRA and guided CBT-based smartphone apps as treatment adjuncts. For caregivers of children and adolescents, weak recommendations were supported for virtual parent meal support training, and moderated online caregiver forums and support groups. For caregivers of emerging adults, guided parental self-help CBT
was strongly recommended, and unguided caregiver psychoeducation self-help was weakly recommended.

**Conclusions:** Several gaps for future work were identified including the impact of sex, gender, race, and socioeconomic status on virtual care among children, adolescents, and emerging adults with eating disorders, as well as research on more intensive services, such as virtual day hospitals.

**Keywords:** Guidelines, Eating disorders, COVID-19, Virtual care, Self-help, Anorexia Nervosa, Bulimia Nervosa, Binge Eating Disorder, Children, Adolescents, Emerging adults
PLAIN ENGLISH SUMMARY

The objective of this project was to develop Canadian Practice Guidelines for the virtual treatment of children, adolescents, and emerging adults with eating disorders. We reviewed the literature for relevant studies, rated the quality of the scientific information within these studies, and then created recommendations for virtual treatments for children and adolescents (<18 years), emerging adults (18-25 years), and their caregivers. We presented our key findings and proposed recommendations to a panel of clinicians, researchers, parents, and those with lived experience from across the country. Based on the discussions by the panel during our presentation, we revised the recommendations accordingly.

This was followed by anonymous voting in an online survey, where panel members could vote and provide comments on the revised recommendations. Our final recommendations include a strong recommendation for in-person medical evaluation when necessary for children, adolescents, and emerging adults; cognitive-behavioural therapy (CBT)-based virtual group therapy, internet CBT-based guided self-help, and internet-based Maudsley Model of Anorexia Nervosa Treatment for Adults (MANTRA) guided self-help for emerging adults; and guided parental self-help CBT for caregivers of emerging adults. For children/adolescents, weak recommendations were determined for telehealth family-based treatment (FBT), online guided parental self-help FBT, virtual parent meal support training, and moderated online caregiver forums and support groups. For emerging adults, weak recommendations were determined for telehealth relapse prevention using MANTRA, guided CBT-based smartphone apps, and unguided caregiver psychoeducation self-help. The panel also identified several areas for future research, including the impact of sex, gender, race, and socioeconomic status on virtual care for eating disorders, as well as the need for further research on more intensive services such as virtual day hospitals.
INTRODUCTION

The negative impact of COVID-19 and the associated social isolation on mental health has been well-described in terms of heightened anxiety and depression (1-3). To cope with the stress of the pandemic, populations around the world are reporting changes in dietary patterns, such as increasing food intake or consuming more comfort food, and feeling anxious due to changed eating habits or activity levels (4, 5). Survey results from a subset of the general population in Italy (n=602), found that almost half of the respondents reported feeling anxious due to changed eating habits (i.e. consuming more comfort food in quarantine, and being inclined to increase food intake to feel better during COVID-19). During this time, females were also more prone to emotional eating than male counterparts (4). In a survey in New Delhi, India, 79.5% of total respondents (n=992) reported that their dietary pattern changed during the country’s lockdown due to stress, especially among people aged 35 to 50 years, who were either increasing or decreasing their caloric intake (relative to before the pandemic) in response to social distancing laws and fears of acquiring COVID-19 (5).

Individuals with eating disorders (EDs) are also experiencing negative impacts related to COVID-19 – a study in Spain found that after only two weeks of quarantine, nearly 40% of adults with EDs reported a worsening of ED symptomatology and almost 60% reported increased anxiety (6). In the USA, Netherlands, and Australia, adults with Anorexia Nervosa (AN) reported increased restriction and fears about not being able to find foods consistent with their meal plan; those with Bulimia Nervosa (BN) and Binge Eating Disorder (BED) reported increases in binge eating and urges to binge; all respondents with EDs noted greater concerns about COVID-19 on their mental health than physical health, as well as concerns of ED relapse related to confinement (7, 8). With closures of ED day hospitals occurring across Canada, Spain, Austria, USA, and the UK, the most severely ill ED patients were left without the intensive treatment they required (6).
There has been a lack of direction in terms of the best options for virtual ED care during the COVID-19 pandemic, and as a result, individuals, families, and clinicians are suffering the repercussions of health care systems that were ill-equipped for such a disruptive event (9). As the ED population is particularly vulnerable and at significant risk of death should they not receive appropriate care (10), there is an urgent need to identify and prioritize the implementation of viable virtual care options. Importantly, evidence indicates that telehealth is a promising alternative to the delivery of outpatient care (11). The telehealth format, however, can present significant challenges for ED care with respect to developing rapport with individuals with EDs, as well as engaging in an appropriate assessment of medical stability. Reports have been made by adults with AN, BN, and BED in the USA and Netherlands about the quality of ED treatment during the pandemic being “somewhat” or “much” worse than usual care (7). While some people with EDs may be willing to engage in virtual therapy, a preference for in-person care remains high (12). Individuals with EDs often experience ambivalence regarding video calls, reporting a heightened self-criticism and awareness of bodily appearance, which respondents indicate creates a negative experience during virtual visits (6). Furthermore, individuals with EDs and caregivers have voiced concern over the lack of face-to-face accountability with virtual weigh-ins, as well as concerns that social isolation may result in long periods of time to engage in excessive exercise (6, 13).

Technologies such as mobile applications (“apps”), self-help resources, and web-based information can be important sources of support during COVID-19 (12, 13). However, social media posts and advertisements continue to bombard users with messages about “inevitable” weight gain related to staying at home, and have been argued to promote shape and weight concerns as well as restrictive and compensatory behaviours among those with, or at-risk for, EDs (14-17). News media reports of food scarcity can cause individuals with EDs to restrict further due to a sense of altruism, or households maintaining a week-long supply of groceries can be a trigger for binge eating episodes (18).
Our team recently published *Canadian Practice Guidelines for the treatment of children and adolescents with EDs* (19); however, many of the recommended interventions are focused on in-person individual and family treatment or group therapy, which currently cannot be delivered. The present project represents an addendum to our guidelines and focuses on the generation of recommendations related to the provision of effective virtual care and online support for children, adolescents, and emerging adults who are living with an ED, and their caregivers. We expect these guidelines to have relevance for clinicians, administrators, and policymakers wishing to provide the best possible ED care during the COVID-19 pandemic and beyond.

**Objectives**

Our aim was to synthesize the best available evidence on: a) the impact of COVID-19 on children, adolescents, and emerging adults with EDs, and their caregivers, and, b) virtual treatments and other online supports and technologies for this population. The research questions listed below were created and discussed by our research team and guideline development panel, consisting of clinicians (e.g. health care professionals), researchers, knowledge users, and patient/parent advocates from across Canada.

**Research questions**

In children/adolescents and emerging adults with EDs and their caregivers: 1) What is the impact of COVID-19? 2) What evidence exists for treatments that can be delivered virtually? 3) What evidence exists for self-help for affected individuals and caregivers? 4) What evidence exists for other e-technology (email therapy, text messaging therapy, smartphone apps)? 5) What evidence exists for virtual day hospital, virtual group therapy, and virtual meal support? 6) When should individuals with
EDs be seen in-person for evaluation? And how can medical monitoring be done at home? 7) How do sex and gender impact virtual care? and, 8) What are the gaps in the research evidence?

METHODS

Overview

Since it was expected that the literature would be limited but diverse, we used scoping review methodology (20-23) to ensure we collated all evidence on the impact of COVID-19 on the ED population, virtual ED treatment in the COVID-19 context, and research focusing on other novel technologies, online self-help support, and individual/caregiver experiences with online approaches. This was followed by a grading of the evidence using the Grading of Recommendations, Assessment, Development, and Evaluation (GRADE) system (24-26). These evidence profiles were then presented to a panel of stakeholders from across Canada, followed by a voting system and arrival at consensus on the recommendations. The Appraisal of Guidelines, Research and Evaluation (AGREE II) tool was used to inform guidelines development and reporting (27).

Virtual care is a broad term which encompasses all the methods in which healthcare providers remotely interact with their patients (28). Telemedicine is one of these methods, as are a variety of video and audio methods that we included in our literature search. The World Health Organization (WHO) defines telemedicine as the delivery of health care services at a distance, by health care professionals using information and communication technologies related to diagnosis, treatment, evaluation, research, or education, all in the interest of advancing health care (29). Generally, the terms ‘telemedicine’ and ‘telehealth’ are synonymous and can be used interchangeably (29). Telehealth can be synchronous, involving the use of audiovisual technology in real-time for patient and health care professional communication, or asynchronous, consisting of health information being collected at one
location and virtually transferred to another for review by a health professional (30). For the purposes of this project, we defined telehealth as meeting over the phone or internet with a therapist in real-time (synchronous). The panel mutually agreed on including telehealth/telemedicine, telephone support services, text messaging, smartphone apps, email, e-mental health platforms (e.g. mental health services delivered via the internet or other digital technologies), self-help, and online forums as types of virtual and online treatment/supports for this study. The panel decided to exclude social media as there was consensus that social media cannot be used to facilitate ED treatment, and should be considered separately. Given the volume of literature that likely exists related to social media and EDs, and the short timeline of our project, it was decided not to include literature on social media and EDs.

**Synthesis methods**

**Eligibility criteria**

Our inclusion criteria were: a) all literature, including quantitative, qualitative, and mixed methods papers on the impact of COVID-19, as well as virtual and online treatments/supports for children/adolescents (<18 years) and emerging adults (18-25 years) with EDs and/or their caregivers; and b) articles written in any language. Our exclusion criteria were: a) studies primarily involving adults (>25 years); and b) studies that did not include EDs or disordered eating behaviours.

**Identifying potentially eligible studies**

**Databases**

A literature search was completed using the following databases: Medline, PsycINFO, EMBASE, Cochrane Database of Systematic Reviews, Cochrane Central Register of Controlled Trials (CENTRAL), and CINAHL. The references of relevant articles obtained were also reviewed.
Literature search strategy

Our library scientist (NB) designed and executed comprehensive searches in the databases listed above to obtain evidence for each of the research questions dating back the last 20 years (2000 to 2020). This time frame was chosen based on the fact that there was likely little or no virtual care prior to 20 years ago and based on feasibility. The searches included a combination of keyword and subject heading for each concept. The sample search strategy included, but was not limited to, various combinations of the following terms as appropriate for the questions being addressed: Anorexia Nervosa OR Bulimia Nervosa OR Eating Disorder Not Otherwise Specified OR eating disorder OR Other Specified Feeding and Eating Disorder OR Avoidant/Restrictive Food Intake Disorder OR Binge Eating Disorder OR Atypical Anorexia Nervosa; AND virtual care OR self-help OR telemedicine OR telehealth OR videoconferencing OR COVID-19 OR coronavirus OR pandemic.

Other strategies

Grey literature was also reviewed, including conference proceedings from the International Conference on Eating Disorders between 2010 to 2020. Databases of ongoing research were searched including CENTRAL and clinicaltrials.gov. We also hand searched the International Journal of Eating Disorders from the last 10 years (2010 to 2020) for relevant articles.

Applying eligibility criteria and extracting data

Two members of the research team independently evaluated the results generated by our searches and came to consensus on which studies met eligibility criteria. We used Endnote and DistillerSR software to organize our studies. Duplicate records were removed. DistillerSR was used for article screening and data extraction. Titles and abstracts were used to exclude obviously irrelevant reports by the two reviewers. Potentially relevant articles were reviewed in full text by two reviewers.
who had to agree on inclusion. Articles in other languages were translated into English using Google Translate (n=6). References of included reviews and book chapters were examined to find other potentially relevant studies. There were no disputes. Data extraction included the indexing of the type of paper, type of control group (if any), methodology, type of virtual intervention, ED diagnosis, age range, sample size, description of intervention, outcomes, results, and whether or not the paper described sex and/or gender as impacting virtual care. Sex was defined as sex assigned at birth, categorized into male or female. Gender was defined as the individual’s self-identified gender role/identity categorized as girl, boy, cis, trans, or other gender identities.

**Appraising studies and guideline-related frameworks**

The *GRADE system* (24-26) explicitly describes how to rate the quality of each study, as well as how to synthesize the evidence and grade the strength of a recommendation. Using this system, we used GRADEpro software to develop an evidence profile for intervention studies where treatment outcomes could be summarized in this fashion. With GRADEpro, we synthesized and classified the overall quality of evidence for each intervention based on the quality of all of the studies, taking into account risk of bias, inconsistency, indirectness, imprecision, publication bias, dose-response, and effect size (25). Although we examined each outcome independently, when the rating of the evidence was the same, we collapsed the treatment outcomes in the GRADEpro tables for the sake of efficiency. We also used the *AGREE II* tool as well as the *Guideline Implementability for Decision Excellence Model (GUIDE-M)* to inform guideline development and this report (31, 32).

**The guideline team**
The Guideline Team is comprised of a core research team and a larger guideline development panel (GDP). This team is skilled in guideline development having just published the first Canadian Practice Guidelines for the treatment of children and adolescents with EDs (19). The research team presented the research questions to the GDP, reviewed evidence summaries, formulated practice recommendations, drafted the guideline, and limited biases that could impeach upon the guideline development process (33-35). The chair of the GDP (MB) is a methodological expert in guideline development, with content expertise outside the field of eating disorders. She led the consensus discussions of the GDP and oversaw conflict-of-interest disclosures and management. Our multi-disciplinary GDP of 27 diverse stakeholders from across Canada included experts in the field of EDs, multi-disciplinary front-line clinicians/knowledge users, those with lived experience (parents and individuals), hospital administrators, and policymakers (all authors on this guideline).

**Procedures**

An initial videoconference (via Zoom) was held on May 29, 2020 with the Guideline Team. The videoconference was recorded to provide an opportunity to review at a later date if necessary. The aims of the videoconference were to confirm the research questions, review the guideline development process and roles and responsibilities of the GDP, and identify potential conflicts of interest. During the meeting, the research questions were refined, the clinical population and outcomes were discussed, and the target audience was reviewed.

Once the literature searches were completed and the evidence profiles generated, a second videoconference (via Zoom) was held on September 25, 2020, which was also recorded so it could be reviewed if required. Two members of the core research team (JC and DP) presented evidence profiles for discussion with the GDP. This was followed by a facilitated discussion of the evidence profiles and
draft recommendations generated by the core team. For each question, the panel reviewed the evidence, and discussed: (i) whether the interpretation of the evidence put forward by the core team aligned with that of the GDP; (ii) strengths and limitations of the evidence base; and (iii) considerations of the generalizability of the studies, precision of the estimates, and whether the evidence aligned with values and preferences of Canadian individuals with EDs and clinicians. Alternative interpretations and suggestions for further research were discussed. Minor or dissenting opinions were noted. Issues regarding implementability of the recommendations were considered, and suggestions for dissemination of the guidelines were elicited. In terms of formulating the recommendations, the panel generally defined a recommendation as ‘weak’ if it was supported by low certainty evidence (e.g. case report, case study, open trial data); a recommendation was defined as ‘strong’ if it was supported by high certainty evidence (e.g. randomized controlled trial data), while also taking into consideration all of the contextual factors mentioned above.

Following the virtual meeting, GDP members were provided with the draft guidelines for review. Group consensus on recommendations and strength of recommendations was obtained by using a modified Delphi method (36), with voting by all team members (except JC, DP, MB and NB) using an anonymous web-based survey platform, Lime Survey (www.limesurvey.com). For a recommendation to be approved, at least 70% of the GDP were required to identify their agreement with the recommendation (26). Consensus was achieved in the first round of voting.

RESULTS

THE IMPACT OF COVID-19 ON EATING DISORDERS

One thousand, three hundred and twenty-five abstracts were identified for review within the IMPACT OF COVID-19 on EDs section of our guideline (see PRISMA flow diagram, Figure 1). After
duplicates were removed, abstracts screened and full-text articles reviewed, 15 papers were included within this section. No additional abstracts were identified through review of reference lists.

**Treatment modifications and/or ED symptoms experienced**

*Children/adolescents.* Some evidence described the impact of COVID-19 on children and adolescents, and/or their caregivers and clinicians. Four descriptive studies revealed a rapid scale-up of telehealth for adolescents in ED programs as a result of the pandemic (37-40). According to clinicians, telehealth created some challenges, including difficulties in ensuring accurate communication when monitoring remotely, privacy concerns, difficulties building rapport with individuals with EDs, individuals with EDs not being physically present for weigh-ins, issues ensuring family involvement in Family-Based Treatment (FBT) sessions, inability to oversee mealtime, and individuals with EDs and/or parents disconnecting or leaving video sessions unexpectedly (38). COVID-19-related modifications to treatment led to an increased workload experienced by clinicians and an increased caregiver burden (e.g. parents having to weigh their child at home) (39, 40). Despite these challenges, in one descriptive study that featured a case report, the individual with an ED (n=1) achieved calorie goals, reduced purging episodes and emotional outbursts, engaged in telehealth care, and avoided re-hospitalization (37).

*Emerging adults.* Like children and adolescents, descriptive study evidence revealed an increase from 0% to 97% in one month for telemedicine visits for emerging adults with EDs in an Adolescent and Young Adult Medicine Clinic and that this population in particular benefitted from telemedicine (41). Challenges included privacy issues and inability to assess anthropometric data (41). If weights and vital signs were unable to be collected at home by caregivers, partnerships with local primary care providers were formed to collect data (41, 42). One qualitative study outlined themes of the impact of COVID-19 treatment modifications for individuals with EDs and caregivers (43). For individuals the themes
included: reduced access to ED services, varying levels of acceptability for remote therapy, reduced motivation for recovery, heightened psychological distress and ED symptoms, and increased attempts at self-management in recovery. For caregivers the themes included: fears of premature discharge from treatment, increased caregiver burden, managing new challenges around patient well-being (e.g. spotting signs of relapse), and new opportunities (e.g. gratitude for increased time at home).

**Impact on clinicians treating individuals with EDs during COVID-19**

*Clinicians caring for individuals with EDs of any age.* Seven articles (44-50) provided suggestions for clinicians for ED care and research during the COVID-19 pandemic, and one article (12) described the impact on clinicians working with individuals with EDs during this time.

Two descriptive studies provided recommendations to combat the challenges associated with remote delivery of ED care (44, 45). These included stressing telehealth sessions were “business as usual” to individuals with EDs who might view telehealth as “second best” using reliable video-call platforms instead of audio calls to deliver virtual treatment, acknowledging the challenges associated with COVID-19 to those with EDs, and having COVID-19-specific ED psychoeducation for CBT or enhanced CBT (CBT-E) sessions. Other descriptive research suggested clinicians consider a different approach to ED treatment during the COVID-19 pandemic, such as implementing a person-centered and harm-reduction approach (46), or specifically for individuals with Avoidant/Restrictive Food Intake Disorder (ARFID), fostering communities of kindness towards ARFID that would involve carefully listening to individuals, family members, and partners of those affected by the disorder and COVID-19 (47). For research, one editorial (49) and one cross-sectional study (50) indicated a need to fast-track ED research and publications related to COVID-19. Additionally, it was recommended that research disruptions be handled by employing technology, reprioritizing study goals (e.g. changing research directions), and encouraging collaboration between sites (49, 50).
In terms of clinician impact, one editorial revealed clinicians’ concern over e-technology adding to their workload, where there may be a new expectation for them to handle patient-related issues during off-work hours given accessiblity to virtual patient data and inquiries (12).

**VIRTUAL CARE AND EATING DISORDERS**

Six thousand, five hundred and fifteen abstracts were identified for review within the virtual care and EDs section of our guideline (see PRISMA flow diagram, Figure 2). Nine additional abstracts were identified through review of reference lists. After duplicates were removed, abstracts screened, and full-text articles reviewed, 69 studies were included within this section of our guideline.

**TELEHEALTH**

**Family-Based Treatment (FBT)**

*Children/adolescents.* Two studies were found using telehealth FBT, which followed an FBT manual but involved therapists virtually assisting parents to support nutritional recovery of their child via a telehealth platform or telephone, in children and adolescents with AN or atypical AN (51, 52).

Significant weight gain from baseline to end of treatment and/or at follow-up was achieved in one open trial (with moderate to large effect sizes; n=10) (51) and in one case report (n=1) (52) (Table 1).

**Relapse prevention (MANTRA)**

*Emerging adults.* In one open trial with individuals with AN, trial completers (n=12) in the Maudsley Model of AN Treatment for Adults (MANTRA) intervention (consisting of 10 sessions of a relapse prevention programme for AN over four months) delivered via telehealth demonstrated increased Body Mass Index (BMI) and lower eating, shape and weight concerns (Eating Disorder Examination – Questionnaire [EDE-Q] scores), compared to baseline (53) (Table 2).

**Cognitive Behavioural Therapy (CBT)**
Children/adolescents. Case report evidence was found for telehealth cognitive and behavioural treatment for ARFID and Other Specified Feeding and Eating Disorder (OSFED); however, both studies had small sample sizes (54, 55). For instance, in one case report, teleconsultations between clinicians and caregivers related to feeding interventions for ARFID resulted in an increase in the frequency of bites of nonpreferred foods consumed, though results were for one individual only (8-year-old male) (54). Likewise, another case report involving CBT telepsychologist visits (using videoconferencing platforms) for Eating Disorder Not Otherwise Specified (EDNOS) resulted in an increase in food intake, improvements in growth, and reduced depression scores; however, again the case report only studied one individual (14-year-old female) (55).

Emerging adults. Two RCTs comparing CBT-based virtual group therapy ("Set your body free" with eight weekly sessions led by a therapist) for emerging adults with probable BN and/or high body dissatisfaction were found (56, 57). One of the RCTs compared face-to-face delivery (n=19) versus internet-delivery (n=21) modes of the program (56) whereas the other RCT compared face-to-face delivery (n=42), internet-delivery (n=37), and delayed treatment control (n=37) (57). Both studies revealed large improvements in body dissatisfaction (Body Satisfaction Questionnaire [BSQ], Body Image Avoidance Questionnaire [BIAQ] scores) and dietary restraint (Dutch Eating Behaviour Questionnaire Restraint Scale [DEBQ-R] scores) in face-to-face and internet-delivery groups. No significant differences between delivery modes were observed in either study (Table 3).

SELF HELP AND GUIDED SELF-HELP

Guided Self-Help

Internet CBT-based guided self-help
Emerging adults. Of all self-help treatments examined, internet CBT-based guided self-help had the most evidence to support its use in emerging adults with AN, BN, BED, and EDNOS. Three RCTs (58-60) and their subsequent studies (61-64) demonstrated reduced ED psychopathology, improvements in ED symptoms (e.g. cessation from binge eating and purging), and/or significant weight gain among various internet CBT-based guided self-help interventions (“Featback” (58), “Overcoming Bulimia Online” (59), and “VIA-Virtual Intervention for AN” (60)) compared to controls. Of these, one RCT that compared intensities of virtual therapist support (low= one time per week versus high= three times per week) in addition to the internet-based program (“Featback”) revealed no added value of therapist support in symptom reduction, but the added support contributed to greater program satisfaction (58) (Table 4).

Non-randomized studies saw mixed results for individuals with BN and/or EDNOS and internet CBT-based guided self-help programs. Two open trials (total n=228) observed significant improvements in ED symptoms (e.g. decreased objective binge eating and vomiting) and general psychopathology following online CBT-based intervention programs (65, 66). One controlled study revealed significant decreases at follow-up in psychopathological levels, binge eating, and vomiting, favouring the intervention group (n=31) over waitlist controls (n=31) (67). In contrast, one case series found that for participants (n =38) who followed an internet CBT-based guided self-help program, there was a notable change in their methods of compensation rather than an actual improvement in behaviours (e.g. participants decreased vomiting frequency but increased excessive exercise episodes) (68). Similarly, one case report with one individual observed no improvements in BN symptoms following a different internet CBT-based intervention (“ProYouth”) (69) (Table 4).

In terms of qualitative findings, two qualitative studies reported that an internet CBT-based guided self-help intervention (“Overcoming Bulimia Online”) was well-received among BN and EDNOS participants (total n=20); these favourable perceptions of the program were attributed to the programs'
accessibility, flexibility, support and content (70, 71). Notably, some participants used the program as a ‘stepping-stone’ to further treatment (70).

**CBT-based Bibliotherapy**

**Emerging adults.** One RCT (72) and its subsequent studies (73, 74) compared internet-guided self-help (INT-GSH) and bibliotherapy-guided self-help (BIB-GSH) among emerging adults with BN and/or EDNOS (n=70 INT-GSH, n=56 BIB-GSH) and reported significant improvements in ED symptoms in both groups. There was no significant difference regarding outcome between delivery modes of the treatment, and authors suggested that both may be equally effective treatment options for this population (Table 5).

**Manual CBT-based guided self-help**

**Emerging adults.** One observational controlled study with individuals with AN studied a manualized CBT-based guided self-help intervention (n=51) with exercises to practice coping skills and improve body image in addition to weekly telephone contact with a clinical psychologist, versus a waitlist control (n=51) (75). Both groups thereafter received inpatient treatment. Duration of inpatient treatment was significantly shorter (by an average of 5.2 days) among intervention participants compared to controls. Body image, slimness ideal, general psychopathology and atypical binges improved significantly during the guided self-help intervention; however, while the intervention group showed more weight gain, changes in BMI did not differ significantly between the groups.

**Internet-based MANTRA guided self-help**

**Emerging adults.** One RCT of individuals with AN studied internet-based MANTRA (iMANTRA) guided self-help (MANTRA workbook and email support from a therapist) in addition to treatment as usual (n=24) versus a treatment as usual only control group (treatment from a local community mental health team or child/adolescent mental health team; n=17) (76). At six months, there was little difference between the groups in terms of outcomes, but at 12 months, the iMANTRA intervention group had a
higher BMI and lower Depression, Anxiety, and Stress Scale (DASS-21) scores than treatment as usual controls; the iMANTRA group also had fewer readmission rates than the treatment as usual control group (Table 6).

**Unguided Self-Help**

*Internet-delivered self-compassionate letter writing unguided self-help*

**Emerging adults.** One RCT with individuals with AN and atypical AN in an internet-delivered self-compassionate letter-writing intervention (adapted from compassion-focused therapy) for non-treatment seeking individuals (n=20) resulted in greater increases in self-compassion and greater decreases in shame and fears of self-compassion, compared to waitlist controls (n=20) (77). The intervention appeared to be acceptable and feasible, but changes in eating pathology (EDE-Q scores) and readiness to get help for one’s weight did not differ between conditions.

**Manual-based Cognitive Remediation Therapy (CRT)**

**Emerging adults.** One qualitative study found that a CRT self-help manual and diary entries for individuals with AN and EDNOS (n=6) were well-received according to high levels of satisfaction and acceptability reported by participants (78). Both individuals with AN and EDNOS and parent participants commented that they would recommend the treatment to others; however, participants suggested that CRT be adapted for delivery via a computer rather than a manual.

**Motivational Enhancement Treatment (MET) and self-help book**

**Emerging adults.** One RCT with individuals with BN and BED compared a self-help only intervention (n=45), consisting of a CBT-based book (“Overcoming Binge Eating”) completed at the participants’ own pace, to in-person MET sessions supplemented with the same CBT-based book (n=45) (79). The MET intervention resulted in increased readiness to change for binge eating and significantly more participants in the MET condition were abstinent from bingeing at follow-up, compared with the self-
help only intervention, but otherwise there were few differences between conditions for eating attitudes and frequency of binge eating and compensatory behaviours. With regard to eating behaviours, participants in both conditions had reduced frequencies of binge eating and compensatory behaviours, but within-group effects indicated that individuals in the MET condition experienced significant reductions in binge eating, compensatory behaviours, and maladaptive attitudes, whereas changes in the self-help only condition were not significant (79).

E-TECHNOLOGY AS ADJUNCTIVE INTERVENTIONS

Moderated online forums

*Children/adolescents.* One qualitative study examining a moderated online discussion forum (n=119 users), which aimed to facilitate support between adolescents regarding their ED and the recovery process, determined several themes illustrating how young people use the forum (80). Themes included taking on the role of the mentor, establishing a safe space online, forming friendships, acquiring help when needed, and seeking peer support for recovery and relapse prevention. The study concluded that moderated online discussions may foster a supportive environment in recovery for children and adolescents with an ED.

*Emerging adults.* One cross-sectional study found that a moderated online forum enabled empowerment for emerging adults experiencing ED symptoms through exchange of information and sharing experiences with others (81). The most pronounced empowering outcome of using the forum was feeling better informed, and to a lesser degree, it increased help-seeking behaviour, optimism, control over the future, confidence in treatment; perceived improvements in the relationships with their therapists. The study determined that forum users had potential to become an active partner in the management of their ED.
Smartphone applications

Children/adolescents. One qualitative study compared the impact of TCAapp between individuals with AN, BN, and EDNOS (n=9), mobile health experts (n=11), health care professionals (n=10), and ED specialists (n=8) (82). TCAapp is a mobile health app that connects children and adolescents with EDs with their therapists in the periods between medical consultation. The study found that most health care professionals considered the app difficult to use, with barriers related to external factors (e.g. lack of time because of workload), while individuals with EDs and ED specialists perceived the app as easy to use. Some individuals with EDs reported barriers related to use of the app including lack of personalization and motivational components, where they also expressed a lack of enthusiasm about the web-based chat option with ED specialists, although this was a facilitator for use from the perspective of ED specialists.

One mixed methods study asked children and adolescents meeting clinical or subclinical criteria for AN, BN, or BED (n=366) about their interest in trying a hypothetical evidence-based mobile mental health app for EDs that included e-coaching (83). Respondents with more severe manifestations of illness were more interested in trying the app, compared to those with less severe signs of an ED. Unwillingness to try the app was related to privacy concerns, worries of parents’ reaction, and feelings that their parents might not want them to participate.

Emerging adults. A variety of evidence was found related to smartphone apps for emerging adults with EDs. One RCT consisted of individuals with AN in a “Recovery Record” plus treatment as usual intervention group (n=20) versus a treatment as usual control group (n=20) (84). Recovery Record is a CBT-based mobile app which involves self-monitoring, encouraging feedback, and coping strategies, with a linking feature with the treating clinician enabling individuals with EDs to share self-entered data with their clinician. At post-intervention, this RCT found non-significant small to moderate between-
group effect sizes favouring the smartphone app intervention group over the treatment as usual group regarding ED psychopathology (EDE-Q scores) and BMI. However, at the 6-month follow-up, there were no significant differences between intervention and control groups for these measures (Table 13).

In terms of non-randomized evidence, one case report saw 108,996 downloads of the Recovery Record app over a two-year period, and of 2503 ratings of acceptability, 84% rated the app as 5/5 (85). From the case report, approximately 50% of Recovery Record app users stated that they do not currently receive ED treatment, suggesting that the app could be effective in reaching an underserved population. One mixed methods study consisting of nine individuals with severe body image and disordered eating concerns determined that a different CBT-based mobile app (“Students Bodies – ED mobile app”) offering one-on-one in-app and phone-based coaching in addition to the app’s core sessions was rated as highly usable by participants (86). The average usability score was originally 78/100, but once modifications were made to the app based on user recommendations this score increased to 83/100. Furthermore, an open trial with 13 universities found encouraging results for the Student Bodies – ED mobile app. Specifically, participant restrictive eating and binge eating significantly decreased over the course of users’ time in the intervention. Vomiting and diet pill/laxative use were not found to significantly decrease over the course of the intervention, but reports of these behaviours were very low (87).

Text messaging

Emerging adults. There was conflicting evidence for text messaging interventions for emerging adults with EDs. In one open trial, 12 individuals with AN, subclinical AN, or BN received personalized, motivational text messages following in-person psychoeducation sessions, which were sent prior to participants’ meal time (88). While the text messaging adjunctive therapy was deemed acceptable and feasible by participants in the open trial, there was no impact on behavioural outcomes including dietary
restraint and kilocalorie intake, and underweight participants reported an increased desire to restrict in response to the text messages. In contrast, a case report with two individuals with BN found that a weekly text-messaging based intervention following discharge from inpatient treatment resulted in positive outcomes, including no binge eating or purging reported over the course of 14 weeks of use (89). The results suggested that the use of text messaging aftercare offers the possibility of supplementing traditional psychotherapeutic treatments.

**Email and/or online counselling**

*Children/adolescents.* Two case reports described individuals with AN (total n=3) that used email as an adjunct to treatment to relay their mood, calorie intake, and ED behaviours to their treating physician (90, 91). In both case reports, individuals generally described email as a positive treatment adjunct, allowing for increased contact with the physician and individuals becoming more aware of their ED behaviours by documenting them online. Weight gain was achieved by one of the two individuals in one case report (16-year-old female) (90). One qualitative study with individuals with any type of ED (n=4) using an online email counselling service reported that the service provided a valuable place for young people to gain additional support and reduce feelings of isolation (92).

*Emerging adults.* Two case reports with individuals with AN or BN (total n=4) found that treatment supplemented with email between the individual and clinician benefitted individuals in terms of improving coping behaviours and creating a greater sense of trust with their therapist (93, 94). Another case report described the effects of a clinician matching two groups of two individuals with AN (total n=4) with similar ED struggles and arranged for them to communicate via email as a means of supporting each other (without involving the clinician) (95). These participants reported positive feelings towards the email matching program, and according to their treating clinician, all progressed well in their treatment. Two qualitative studies with individuals with AN, BN and EDNOS (total n=309)
found email online counselling was rated positively amongst users, with the supportive comments, fast and easy contact, counsellors’ competence related to EDs, and the service being free of charge as some of the main facilitators for using the online service (96, 97). Two cross-sectional studies (98, 99) and one mixed methods study (100) were related to the same online consulting service, which involved anonymous emails with a health care professional for free via a website. All three studies found that the service was often the initial point of contact between users with AN, BN, and BED and a professional to get help for their ED; relatives of those with EDs were also accessing the online consulting service to learn more about EDs (98-100). Finally, one open trial with individuals with BN, BED, and EDNOS (n=23) further described the impact of using email to communicate food intake and symptoms with a therapist and in return, receiving CBT-based or eclectic support from the therapist (101). Results indicated significant improvements in depressive and bulimic symptoms at 3-month follow-up (101).

CAREGIVER INTERVENTIONS FOCUSED ON CHILD OUTCOMES

Online guided parental self-help – FBT

Children/adolescents. One case series with 19 families found that the individuals with AN in the online guided parental self-help FBT intervention experienced weight gain similar to clinician-delivered FBT programs, with improvements in ED-related psychopathology (EDE-Q scores) also reported by the end of the treatment (102). Similarly, after participating in online family-based early intervention pre-recorded sessions, individuals in one open trial (n=12 diagnosed with AN, n=12 at risk for AN, and n=22 at high-risk for AN) remained stable or increased in ideal body weight by post-intervention (103) (Table 7).

CAREGIVER INTERVENTIONS FOCUSED ON CAREGIVER OUTCOMES

Unguided caregiver self-help using web-based and/or pre-recorded videos focused on psychoeducation and communication skills
Children/adolescents. One mixed methods study that involved a Meal Support Training program (pre-recorded videos and an accompanying manual) was well-received by families with a child with AN, BN or EDNOS (n=40) (104). Caregivers reported that the program was informative, convenient, and useful in fostering caregiver understanding and patience with their child and the ED recovery process (Table 8).

Emerging adults. Two RCTs had mixed evidence for effectiveness measured by caregiver outcomes (105, 106). One RCT that compared a web-based group (n=23) and an in-person workshop group (n=27) for caregivers of emerging adults with AN and BN revealed improvements in the Caregiver Accommodation and Enabling Scale for EDs favouring the web-based intervention, while changes in caregiver burden favoured the workshop (105). Another RCT with caregivers of emerging adults with AN, BN, atypical AN, atypical BN, and EDNOS that compared a DVD intervention (n=147) to a control condition (n=138) found Caregiver Accommodation and Enabling Scales for EDs were not reduced by the DVD intervention, but caregiver burden was reduced by the intervention compared to controls (106) (Table 9).

Guided caregiver self-help – Skills

Children/adolescents. One mixed methods study, consisting of 16 caregivers for individuals with AN or BN watching DVDs on ED care with supplemental telephone coaching, found caregiver general distress (measured by General Health Questionnaires scores) decreased significantly from baseline to post-intervention (107). Caregivers also expressed high levels of satisfaction with most aspects of the intervention and reported improvements in psychological distress and depression following the intervention, although these measures did not reach statistical significance (Table 10).

Guided parental self-help CBT (skills training approach + workbook or CBT-based online modules)

Emerging adults. Three RCTs studying variations in guided caregiver self-help based in CBT reported somewhat similar findings in terms of expressed emotion and ED symptom impact scale for caregivers of emerging adults with AN (76, 108, 109). One RCT found its skills training intervention group
(“Experienced Carers Helping Others” [ECHO]; n=134) had reduced caregiver expressed emotion levels and ED symptom impact scale scores, compared to the control group (n=134) (76). Another RCT found a CBT-based online module intervention (“Overcoming Anorexia Online”; n=33) significantly reduced caregivers’ anxiety and depression at post-treatment compared to a telephone and email hotline support control group (n=30) (108). Other outcomes, including caregivers’ expressed emotion and ED symptom impact scores, also had greater reductions in the intervention group than the control group, however these were not statistically significant (108). The third RCT compared a CBT-based online module intervention (“Overcoming Anorexia Online”) plus professional guidance via telephone (n=19) to a group that received the “Overcoming Anorexia Online” intervention alone (n=18), and reported similar reductions in caregiver expression of emotion and ED symptom impact scale scores between both groups, but no significant difference between groups (109) (Table 11).

**Moderated online caregiver forums**

**Children/adolescents.** Two non-randomized studies evaluated the use and impact of moderated online forums for caregivers of children with EDs, both describing online tools positively (110, 111). In a qualitative study, five mothers explained that they use moderated blogs (available on the FEAST-Families Empowered and Supporting Treatment of Eating Disorders website) as a tool to foster social support alongside FBT sessions (110). One open trial that consisted of 13 caregivers with a child with AN engaging in weekly therapist-guided virtual chat sessions had high satisfaction ratings (91.7%) in addition to reports that the sessions were accessible, convenient, and easy to use (111). Caregivers also reported looking forward to the chat sessions as they assisted in coping with their child’s ED, and that they would recommend the chat to others (111) (Table 12).

**Children/adolescents and emerging adults.** A case report using teleconferencing for caregiver (n=6 parents) support for any ED in youth between the ages of 12 and 22 found this approach may be
promising for developing healthy caregiver coping mechanisms during their child’s recovery (112).

However, these authors concluded that in-person meetings may be more ideal (Table 12).

IN-PERSON CARE VERSUS HOME MONITORING

In-person medical evaluation

Emerging adults. Some evidence was found describing factors to consider when deciding if individuals with EDs should be seen in-person for evaluation following their remote care in accordance to COVID-19 social distancing regulations. A descriptive study indicated that a person with an ED should be asked to come in to a clinic for limited in-person sessions if they were clearly engaging in concerning behaviours (e.g. increased restricting, purging, over exercising) according to self or caregiver reports or if the provider or caregiver had a high index of suspicion for medical deterioration (41). Furthermore, a commentary described that in the COVID-19 context, ED programs utilized weight and heart rate criteria to define urgency and need for in-person medical assessments, either from temporary “vital signs clinics” where individuals with EDs would come in to a clinic for daily heart rate, blood pressure, and weight assessment (that was either followed by or preceded by a telehealth visit), or some programs enlisted parents to monitor and report vital signs including teaching parents to take pulses and assessing weights on home scales (42).

Home monitoring

Children/adolescents. There was some limited evidence from an open trial of the effectiveness of wearable sensors for children and adolescents with AN (n=27) to monitor heart rate and heart rate variability by a clinician (not by caregivers) in a remote setting (113) (Table 14). However, other recent evidence described interventions for home monitoring to be performed by caregivers of children with EDs during the COVID-19 pandemic. For example, two descriptive studies discussed home monitoring
for children and adolescents with AN in which caregivers were enlisted to measure their child’s weight, heart rate, and blood pressure (Table 14), as well as record food intake and estimate energy expenditure, and report all findings to the clinician during telehealth sessions; clinicians were to visually inspect individuals with EDs during telehealth visits (37, 40). Overall, these new responsibilities for caregivers may contribute to a higher caregiver burden. Other research suggests that only weight be monitored by caregivers at home (38).

SEX, GENDER AND OTHER CONSIDERATIONS

There was no evidence found related to the impact of sex, gender, and other considerations on virtual care for children/adolescents or emerging adults.
RECOMMENDATIONS

Section 1. Telehealth

Telehealth FBT may be a reasonable treatment option for children and adolescents with AN.

Weak Recommendation.

Qualifying Statements:
Family-Based Treatment (FBT) focuses on empowering parents to re-nourish their children. Although the evidence is rather scant, the panel member clinicians indicated that due to COVID-19 many of them are now using FBT by telehealth in their routine clinical practice and they suggest that there is good clinical reason to support these methods. Although these are treatment adaptations to a virtual mode of delivery, this treatment is known to be effective in-person, which adds confidence that it is likely beneficial when delivered virtually as well. There is urgency to adopt these treatments and likely minimal harm involved in delivering them by telehealth, as long as individuals are properly monitored medically. Some clinicians have been reluctant to have parents weigh their own children which would be a necessary component of FBT by telehealth.

Key Evidence:

- Significant weight gain from baseline to end of treatment and/or at follow-up was seen among individuals with AN and atypical AN in one open trial (with moderate to large effect sizes; n=10) (51) and in one case report (n=1)(52).
Telehealth relapse prevention using MANTRA may be a reasonable treatment option for *emerging adults* with AN.

**Weak Recommendation.**

**Qualifying Statements:**
Maudsley Model of AN Treatment for Adults (MANTRA) aims to address the cognitive, emotional, relational and biological factors which tend to maintain AN. This study followed inpatient admission/day patient treatment for AN. Eight sessions were delivered by videoconference; the first and last sessions were in person. Although this was an open trial, this intervention could be quite useful for those leaving intensive treatment and could provide a bridge to outpatient care.

**Key Evidence:**
- At the end of treatment (4 months, 10 sessions), an open trial with individuals with AN (n=16) resulted in increased BMI and reduced eating, shape, and weight concerns (EDE-Q scores) among participants (53).

CBT-based virtual group therapy is strongly recommended for *emerging adults* with BN and/or those with high body dissatisfaction.

**Strong Recommendation.**

**Qualifying Statements:**
Studies used a cognitive behavioural program called “Set your body free”. This program included eight weekly 90 min group therapy sessions by videoconference (6-8 participants per group). Participants had
high degrees of body dissatisfaction and the authors indicate that they had probable BN, but it is difficult
to make firm conclusions as a diagnosis of BN was not required to participate in the sessions. Our
recommendation suggests that those with BN would likely benefit. One issue with virtually delivered
treatments is that the generation of a diagnosis is more challenging and sometimes a diagnostic
interview is not completed.

**Key Evidence:**

- Two RCTs with those exhibiting BSQ scores above the community mean (>81.5) and/or those
who possibly had BN (according to BULIT-R scores): n=19 face-to-face delivery and n=21
internet-delivery (56); n=42 face-to-face delivery, n=37 internet-delivery, and n=37 delayed
treatment control (57). Both face-to-face and internet groups showed large improvements in
body dissatisfaction (BSQ, BIAQ scores) and dietary restraint (DEBQ-R) (compared to control in
the RCT with a delayed treatment control group); no significant differences between face-to-
face and internet delivery modes were observed (56, 57); both were effective.

**Additional Promising Interventions**

- Telehealth cognitive and behavioural treatments for ARFID and OSFED require more study (54,
55). Although there is currently no evidence for family interventions for OSFED and ARFID
delivered by telehealth, this should also be a focus of research.
Internet CBT-based guided self-help is strongly recommended for emerging adults with AN, BN, BED, and EDNOS, as well as relapse prevention in AN.

**Strong Recommendation.**

**Qualifying Statements:**

There is strong evidence that internet CBT-based guided self-help is effective for emerging adults with AN, BN, BED, and EDNOS, and possibly for relapse prevention in AN. These young adults likely need to be motivated for recovery in order to benefit. Although CBT-based bibliotherapy had slightly less evidence (compared to CBT-based internet therapy), there was still some evidence that it was beneficial. The panel emphasized that many parents request psychoeducation on EDs, so there was endorsement of bibliotherapy not only for individuals but for parents as well. The panel recognized that books and workbooks on EDs may be especially useful if no internet or computer is available. One pitfall to all of these treatments is that speaking and reading in English is required, since all of the treatments included in this current review required speaking and reading in English, but it should be noted that translations are becoming available for some books on EDs.

**Key Evidence:**

CBT – Internet Guided Self-Help

- One RCT with those self-reporting AN, BN, BED, or EDNOS symptoms, assessed by the Short Evaluation of Eating Disorders (SEED): total n=87 Featback, n=88 Featback + low-intensity therapist support, n=89 Featback + high-intensity therapist support; n=90 waitlist control. Interventions were superior to control in reducing ED psychopathology (SEED and EDE-Q scores). No added value of therapist support in symptom reduction but contributed to greater
satisfaction. No significant differences between therapist support conditions. Lowest costs in condition with low-intensity therapist support (58).

- One RCT with those diagnosed with BN or EDNOS: n=38 “Overcoming Bulimia Online” intervention; n=38 waitlist control. Intervention group had higher rates of cessation from binge eating and purging than delayed treatment condition; gains maintained or continued to improve at follow-up (59).

- Relapse prevention in AN – one RCT: VIA intervention (relapse prevention based in CBT after AN inpatient treatment) n=128, control n=130. Moderate dropout rate (15.5%); intervention completers gained significantly more weight (individuals with AN) than controls; favourable course for BMI-adherence to intervention, more spontaneity, better self-esteem (60).

CBT-based Bibliotherapy

- One RCT with those diagnosed with BN or EDNOS (with bulimic symptoms): total n=70 internet guided self-help (INT-GSH); n=56 bibliotherapy guided self-help (BIB-GSH). ED symptoms (binge eating and compensatory behaviour) improved significantly in both groups (no significant differences regarding outcomes between groups). No group differences in Eating Disorder Inventory (EDI) subscales (adolescents n=29, adults=97) (72).

Internet-based MANTRA guided self-help is strongly recommended for emerging adults with AN.

Strong Recommendation.

Qualifying Statements:
Internet-based Maudsley Model of AN Treatment for Adults (iMANTRA) involved a workbook and email support by a therapist 1-3 times a week for the first 6 months, then 1 time per week for months 7-12. Workbook content included nutrition planning, coping strategies, and strategies to reduce anxiety.

**Key Evidence:**

- One RCT with individuals with AN n=24 iMANTRA intervention + treatment as usual; n=17 treatment as usual. At 6 months, there was little difference between groups; at 12 months, the intervention group had a higher BMI and lower scores on Depression, Anxiety, and Stress Scale (DASS-21) than controls (76).

**Additional Promising Interventions**

- Internet-delivered self-compassionate letter writing unguided self-help for *emerging adults* with AN and atypical AN requires more study (77).

**Section 3. E-technology as Adjunctive Interventions**

Guided CBT-based smartphone apps may be reasonable adjunctive treatment options for *emerging adults* with AN.

Weak Recommendation.

**Qualifying Statements:**

There was much discussion on the panel regarding the CBT app (Recovery Record) for emerging adults. The context in which the study was done was post inpatient admission for AN in emerging adults for a
period of 8 weeks with feedback from a therapist on a weekly basis at minimum. Some panel members felt we should ensure the app was recommended only as an adjunct to standard treatment, offered concurrently. Some wondered if there could be unintended harms by individuals with EDs believing that they were in treatment by use of the app, when in fact they were not really making progress toward recovery, or that the app did not provide the level of treatment needed for those with more severe EDs. Some felt that these apps could serve a purpose in meeting young people where they are at in terms of level of motivation and that such an app might set them on a pathway to connect with more standard care. The app might also be able to reach more people who could not access standard care. We agreed to leave this as a weak recommendation, with the caveat that in more severe presentations of EDs, the app may be insufficient, and that the app should be used as it was in the study with guidance from a therapist. In fact, the panel felt it should only be endorsed as an adjunctive treatment, accompanying standard treatment.

**Key Evidence:**

- One RCT with emerging adults diagnosed with AN: at post-intervention, non-significant small to moderate between-group effect sizes favoured the smartphone app intervention group with therapist feedback (received support from therapist through the app two times per week for 25 minutes each for the first four weeks, which decreased to once per week during the last four weeks; n= 20) regarding ED psychopathology (EDE-Q scores) and BMI vs. control (n= 20); at 6-month follow-up: no significant differences between intervention and control groups for these measures (84).

**Additional Promising Treatments**
• Email therapy (as a treatment adjunct) for children and adolescents with AN requires more study (91).

• Email therapy and/or online counselling (as a treatment adjunct) for emerging adults with AN, BN, BED, and EDNOS requires more study (93, 94, 100, 101).

• Text messaging therapy (as a treatment adjunct) for emerging adults with AN, subclinical AN, and BN requires more study (88, 89).

Section 4. Caregiver Interventions Focused on Child Outcomes

Online guided parental self-help FBT may be a reasonable treatment option for children and adolescents with AN, at risk for AN, and at high risk for AN.

Weak Recommendation.

Qualifying Statements:

Research evidence evaluates the use of pre-recorded videos that parents can access via a web-based platform, along with coaching by a therapist via phone or videoconferencing. The videos focus on the principles of FBT including urgency to act, caregiver empowerment, externalization of the ED, agnosticism, and psychoeducation around medical complications. One study involved children and adolescents diagnosed with AN (according to DSM-5 criteria for AN), and the other included those with AN, at risk for AN, and at high risk for AN, as determined by a diagnosis of AN within the past six months and/or screening results using established risk factors for EDs, retrospective correlates, and/or early symptoms of AN.
**Key Evidence:**

Online guided parental self-help – FBT

- One case series: n=19 families. At the end of the treatment and at follow-up, individuals with AN experienced weight gain similar to standard clinician-delivered FBT programs. ED-related psychopathology (EDE-Q scores) of the children improved by end of treatment (102).

- After the family-based early intervention sessions (six online sessions), individuals in 1 open trial (n=12 diagnosed with AN, n= 12 at risk for AN, and n=22 at high risk for AN) remained stable or increased in ideal body weight (103).

**Section 5. Caregiver Interventions Focused on Caregiver Outcomes**

Virtual parent meal support training may be beneficial for caregivers of children and adolescents with AN, BN, and OSFED.

Weak Recommendation.

**Qualifying Statements:**

Caregivers watched pre-recorded video content and receive a manual on meal support strategies.

**Key Evidence:**

- One mixed methods study: the virtual Meal Support Training program was well-received by families (n=40; indicated by the lowest rating of the program being 3.79 [SD=0.843] on a 5-point scale [1=not informative, 3= somewhat informative, 5=very informative]), and families deemed
the intervention as informative and convenient; caregivers reported that the program helped them to be more understanding and patient with their child and the ED recovery process (104).

Unguided caregiver self-help using web-based and/or pre-recorded videos focused on psychoeducation and communication skills, may be beneficial for caregivers of emerging adults with AN, atypical AN, BN, atypical BN, and EDNOS.

Weak Recommendation.

**Qualifying Statements:**

One study focused on a web-based intervention with 8 modules and a workbook. Core elements of the intervention were ED psychoeducation, communication skills, meal support strategies, caregiver needs. The other study provided 5 pre-recorded information sessions to caregivers on the impact of EDs, meal support strategies, communication skills, motivational stages of change. There was no contact with a therapist.

**Key Evidence:**

- One RCT with caregivers for individuals with AN and BN: n=23 web-based group; n=27 workshop group. Improvements in Caregiver Accommodation and Enabling Scale for EDs favored web-based intervention; changes in caregiver burden favored workshop intervention (105).
- One RCT with caregivers for individuals with AN, BN, atypical AN, atypical BN, and EDNOS: n=147 DVD intervention; n=138 control. Caregivers’ Accommodation and Enabling Scale for EDs scores were not reduced by DVD intervention, but caregiver burden was reduced by intervention compared to control (106).
Guided parental self-help CBT (skills training approach + workbook or CBT-based online modules) is strongly recommended for caregivers of emerging adults with AN, and may be effective for caregivers of children and adolescents with AN or BN.

Strong Recommendation.

Qualifying Statements:
One study used Experienced Carers Helping Others (ECHO)-guided self-help skills training for caregivers using a book, pre-recorded videos and telephone coaching sessions. Another two studies used a web intervention using interactive multimedia CBT for caregivers to help them understand and meet their own needs; the latter intervention was called Overcoming Anorexia Online (OAO). The study involving caregivers of children and adolescents used a series of pre-recorded videos on communication skills, cycle of change, and strategies for supporting eating. Telephone or email support was also provided by a clinician. The panel agreed that these interventions for caregivers should be used in addition to treatment for the individual with an ED and should not be confused with actual treatment.

Key Evidence:
Guided self-help for caregivers of emerging adults

- One RCT with caregivers for individuals with AN: skills training intervention (ECHO) (n=134) had reduced caregivers’ expressed emotion levels and ED symptom impact scale scores, compared to controls (n=134) (76).

- One RCT with caregivers for individuals with AN: Compared with telephone and email hotline support control (n=30), the CBT-based online modules intervention group (OAO; n=33)
significantly reduced anxiety and depression in caregivers at post-treatment; caregivers in the
OAO intervention group also had greater reductions in expressed emotion and ED symptom
impact scale scores compared to controls, but these were not significant (108).

- One RCT with caregivers for individuals with AN: Caregiver expressed emotion and ED symptom
impact scale scores were similarly reduced in both groups (n=19 OAO + guidance; n=18 OAO
only); no significant difference between groups (109).

Guided caregiver self-help – Skills for caregivers of children and adolescents

- One mixed methods study with caregiver for individuals with AN and BN (n=16). From baseline
to post-intervention, caregiver general distress (General Health Questionnaire scores) decreased
significantly (107).

Moderated online caregiver forums and support groups may be beneficial for caregivers of children
and adolescents with AN.

Weak Recommendation.

Qualifying Statements:

Although the evidence was minimal, the online forum for caregivers was given a weak recommendation
due to the enormous benefit to parents, and well as popularity of the Maudsley Parents forum. A note
to readers here is that we did not include the search term ‘social media’ in our database searches; the
panel agreed a priori that the research related to social media would be too large to review and would
not meet our goals of answering our research questions related to the best treatments that can be
delivered virtually or in the COVID-19 context. Thus, we focused on moderated forums (as opposed to
unmoderated forums). The panel mentioned the importance of parent support as helpful in reducing caregiver burden. From their own experience running virtual parent-led support groups, many parents would prefer in-person support, but are glad there is a virtual option currently being offered in some locations. Virtual options might have benefits in terms of time and being able to access from home (no need for caregivers to arrange childcare).

**Key Evidence:**

- One open trial with caregivers for individuals with AN: virtual chat room sessions were highly feasible and acceptable (91.7% satisfaction rating) among caregivers (n=13), who reported sessions as accessible, convenient, and easy to use (111).
- One case series of teleconferencing for caregiver support for caregivers of those with any ED (not specified) aged 12-22 years found benefit for caregivers (112).

**Section 6. In Person Care versus Home Monitoring**

It is strongly recommended that individuals (of all ages) with EDs are seen in-person for medical evaluation by specialists in conjunction with local care providers when necessary and that international guidelines and criteria for admission are followed.

**Strong Recommendation.**

**Qualifying Statements:**

This recommendation is as based on expert consensus. The panel was unanimous in their agreement that it is not possible to offer all care virtually to this patient population. Guidelines by the Society for
Adolescent Health and Medicine provide criteria for hospital admission and medical monitoring (114) in children, adolescents, and emerging adults and should be followed by all practitioners. Individuals with eating disorders must be able to access emergency rooms and hospital beds despite competing demands for resources in the COVID-19 era. The threshold for admission to hospital should not be changed due to the COVID-19 context. In fact, a direct relationship may exist between the pandemic and an increase in the number of young people needing admission to hospital for EDs. This was mentioned by panel members and there is preliminary evidence to support this (personal communication with several pediatricians across the country, and a study from Perth, Australia (115)).

In terms of monitoring at home, weight can be taken at home on a home scale, however FBT therapists should educate caregivers around possible falsification of weight. Monitoring of vital signs at home is not recommended due to difficulties in proper measurement interpretation and issues with methods of monitoring (may not be reliable or accurate) that could cause harm. Vital signs should be monitored by a health professional. Lack of progression in virtual therapy, concerns around accuracy of home weights, or new symptoms should signal the need for in-person evaluation. Additional reasons for necessary in-person medical evaluation may include lack of reliable measurements at home, caregiver or patient unable or unwilling to monitor weight at home, and the patient’s situation being medically precarious.

Suicidality is another reason individuals may need to be assessed in-person. Should a young person not wish to come to hospital, but there is grave concern over physical or mental health, legal processes should be followed in order to ensure that the individual receives the assessment and treatment that they require.

Key Evidence:
• Some limited evidence of the effectiveness of wearable sensors for adolescents with AN to monitor vital signs by a clinician, not by caregivers (n=27) (113).

• Some studies indicate that with rapid scale up of virtual FBT, caregivers for those with AN were given the role of monitoring weight at home (37, 40).

Section 7. Sex, Gender and Other Considerations

We strongly recommend that equity-seeking groups and marginalized youth should be provided equal access to treatment.

Strong Recommendation.

Qualifying Statements:
This recommendation was not based on research evidence, but rather on expert consensus. Equity-seeking groups as defined by the Public Service Alliance of Canada include racialized people, people with disabilities, Aboriginal (First Nations, Inuit and Métis), women, lesbian, gay, bi-sexual, and trans individuals (116). Trans-youth and boys are at high risk of not being recognized by providers as having eating disorders. Providers need to be more vigilant with these groups, and also be aware of the barriers to care that they face. In addition, non-English speaking youth and their families struggle with extra barriers to care, especially in a virtual world. It is difficult for an interpreter to be involved virtually, although this should be attempted. Those in rural areas or of lower socioeconomic status may not have access to the internet, or they may face limitations related to internet speed and high cost. Providers should also be vigilant to violence occurring in homes as individuals experience heightened isolation. Health care workers should be aware of how racism affects the quality of and access to health
care for racialized groups and should seek to reduce barriers. Indigenous peoples, especially in the far north, face additional barriers to care related to geographical isolation, lack of access to the internet or reliable internet, lack of access to videoconferencing centres, as well as lack of confidentiality due to small population sizes and therefore, increased likelihood of kinship, peer, or social connections to health care providers. Inclusion of equity-seeking groups in research is essential.

**Key Evidence:**

No evidence could be found in our search on these topics.

**DISCUSSION**

These are the first Canadian Practice Guidelines to evaluate the evidence on virtual-based treatment focused specifically on children and adolescents (<18 years) and emerging adults (18-25 years) with EDs, in the COVID-19 context. Strong recommendations were supported in favour of in-person medical evaluation when necessary for children, adolescents, and emerging adults with EDs. In addition, for emerging adults, CBT-based virtual group therapy for BN, internet CBT-based guided self-help for AN, BN, BED, and EDNOS, and internet-based MANTRA guided self-help for AN were strongly recommended. Weak recommendations were generated for telehealth relapse prevention using MANTRA for AN, and guided CBT-based smartphone apps as treatment adjuncts for AN. For children and adolescents with AN, weak recommendations were supported for telehealth FBT, and online guided parental self-help FBT. In terms of caregiver outcomes, guided parental self-help CBT for caregivers of emerging adults with AN or BN was strongly recommended, while unguided caregiver psychoeducation self-help was weakly recommended for most ED diagnoses. For caregivers of children and adolescents...
with EDs, weak recommendations were supported for virtual parent meal support training, as well as moderated online caregiver forums and support groups.

Panel members generally commented on the pros and cons of virtual care. While virtually delivered care may be easily accessible for those with a reliable internet connection at home, this poses a potential barrier to care for families without home internet access. Clinicians mentioned fatigue from virtual care and challenges navigating the legal and procedural aspects of technology-based care. Others mentioned that their patients and families often do not like to see their own image, which poses challenges when trying to conduct video call appointments. Suggestions for this problem were proposed on the panel including having the family blocking their own image by covering this portion of their own screen (e.g. hiding their image on the videoconferencing platform or using a sticky note on their monitor). Some panel members also indicated it is hard to evaluate weight status in patients by virtual means. This resonates with the literature already published on this topic. Three studies identified ED professionals’ opinions on the impact of digital technologies on their patients, indicating potential benefits and drawbacks. Specifically, in two qualitative studies (total n=65) professionals described advantages in virtual care including better patient-clinician communication and access to patient reported data (via apps) ahead of scheduled sessions. However, disadvantages included added workload for the clinician and the risk that it may be easier for patients to continue with and hide their ED symptoms and behaviours in an online environment (117, 118). Survey results from one cross-sectional study found that professionals (n=282) believe e-health may be a more useful treatment for adolescents and adults than children and seniors, and may be more beneficial for anxiety and depressive disorders than for addiction and eating disorders (119).

An additional aspect to consider is how to appropriately train clinicians in virtual modalities. Although thought to be equally efficacious, virtual adaptations of treatments may have differences that
could affect efficacy. Three studies observed the effectiveness of various web-centered ED training programs for health care professionals. In one RCT that compared a CBT-E web-centered supported training group (receiving phone call support in addition to the course components; n=81) to a CBT-E web-centered independent training group (only received access to the CBT-E course; n=75), it was confirmed that web-centered training can successfully train therapists to deliver CBT-E (120). The supported and independent training programs were also equally effective as both training groups had increased scores in therapist competence from pre-intervention to post-intervention (120). This CBT-E web-centered training program was also studied in an open trial (n=765), similarly confirming that competence in delivering CBT-E was improved following web-based training. Factors associated with a beneficial effect from this training included compliance with training recommendations, higher number of training modules completed, and treating at least one patient while in training (121). In a separate open trial that involved a tele-mentoring project between ED specialists (mentors) and community-based practitioners (mentees) (Project ECHO; n=99) which aimed to disseminate specialized ED knowledge to clinicians in need and located in underserved areas via Zoom, tele-mentoring was found to be promising to bridge the speciality-based knowledge gap between ED-trained and front-line clinicians (122).

Panel members also discussed the various platforms available for virtual care delivery. There is limited evidence describing what online platform is the most ideal to deliver ED care virtually, and as such, clinicians tend to only use what is available through their institution. However, one descriptive study that summarized the opinions of clinicians delivering CBT for EDs via telehealth during COVID-19 revealed positive experiences about working with cloud-based videoconferencing and live-chat services, including Zoom, Facetalk, Google Meets, Vsee, and Microsoft Teams (44). Other software choices, such as Skype and FaceTime were seen as less reliable and less secure in comparison, and therefore were not
often recommended (44). According to this descriptive study, platforms such as Zoom that allow for confidential sharing of documents during sessions, high security, white board feature, and ease of use may be the most favourable to support virtual care. However, other platforms endorsed by Canadian health care agencies as meeting privacy and security requirements (e.g. Skype for Business and Zoom Healthcare) were not evaluated in this particular study. Our panel also did not endorse one single platform; panel members identified benefits to several different platforms and acknowledge that practitioner context and organizational mandates may be a key factor in the selection of virtual technology.

Two studies described general recommendations for clinicians delivering virtual mental health interventions during COVID-19. One commentary focused on recommendations for improving the delivery of virtual mental health care, such as: programs offering training and supervision in virtual therapy tools, national licenses being implemented to practice virtual therapy (telehealth), providers being taught to ensure patient confidentially during virtual sessions, broadening reimbursement coverage to include evidence-based virtual therapies, continuous evaluation of the efficacy of virtual therapies on broader mental health disorders, and introducing innovative and timely virtual mental health practices into health care systems (123). One descriptive review noted that guidelines for telepsychiatry in EDs are lacking, but in general, clinicians involved in telepsychiatry during COVID-19 should: a) familiarize themselves with the telehealth system to ensure sessions run smoothly; b) focus on effective communication during sessions; and c) document any issues with telehealth immediately, so that improvements can be made for future sessions (124).

Strengths
The strengths of this guideline are numerous. We used a rigorous and evidence-based methodology for our scoping review and our guideline development. Our scoping review methods included a thorough review of all literature (including a variety of databases, grey literature, and papers in any language), and we had few exclusion criteria. We translated six papers in German or French into English for full-text review and we also examined the references of included reviews and book chapters to ensure we did not miss any relevant studies. In terms of guideline development, conflict of interest statements were reviewed by an impartial chair (MB) in order to address any biases. We had an initial virtual meeting to ensure that the research questions were unanimously agreed upon by the panel, as well as a second virtual meeting to discuss our recommendations, which was followed by an anonymous voting procedure. Furthermore, our panel included the voices of various stakeholder groups including researchers, clinicians, policymakers, parents, and those with lived experience of receiving treatment for an ED.

Limitations

Although thorough, our search strategy had limitations. We were unable to retrieve seven citations as full text articles as they could not be located. We also attempted to examine sex differences, but the numbers of male subjects were so small that no conclusions could be drawn, and studies did not comment on the impact of sex on virtual care. While we searched the literature thoroughly for virtual day hospital and virtual meal support, we were unsuccessful in finding any articles on these topics. We found one mixed methods study related to virtual meal support training (pre-recorded videos) for caregivers. Furthermore, virtual care presents some difficulties related to diagnostic certainty. Many studies enrolled participants based on symptom evaluation using instruments rather than diagnostic interview, with study inclusion criteria also bridging DSM-IV and
DSM-5 criteria. Therefore, recommendations for specific ED diagnoses may be seen as suggestive rather than definitive. Despite these limitations, these guidelines represent a significant step forward in adopting virtual care in the field of ED for treatment among treatment-seeking children/adolescents and emerging adults and their families.

**Future directions**

Several gaps were noted by the guideline panel, which should be a focus for future study. These gaps include the impact of sex, gender, and underserved populations on virtual care among children/adolescents and emerging adults with EDs, as well as the efficacy of more intensive treatments such as virtual day hospital, or virtual meal support therapy. Mixed methods studies may be a useful study design for these understudied areas, to understand both quantitative and qualitative impacts.

There were several other gaps identified by the panel as needing further study. The panel was unable to make any recommendations for adjunctive email therapy and/or online counselling, and text messaging therapy. While some encouraging evidence was found in these areas for children/adolescents and emerging adults with AN, BN, BED, and EDNOS, the panel concluded that these therapies are promising in terms of possible adjuncts to treatment, however, should not be recommended without further study. The panel suggested that creative interventions be developed for those on a waiting list or transitioning in levels of care. Such a possibility might be self-compassionate letter writing unguided self-help (adapted from compassion-focused therapy) for emerging adults, (77) but again more research is required to support this recommendation. Furthermore, there was some positive evidence for telehealth cognitive and behavioural treatment for children/adolescents with ARFID and OSFED, but additional research is also required to draw conclusions, as sample sizes in these studies were small.
Finally, research on telehealth family interventions for ARFID and OSFED is lacking but should be a focus of further study.

CONCLUSIONS

For children/adolescents and emerging adults with EDs, our Canadian Practice Guidelines for virtual treatment during COVID-19 recommend the provision of: 1) in-person medical evaluation when necessary (strong recommendation). For children/adolescents with AN, these guidelines recommend: 1) telehealth FBT (weak recommendation) and 2) online guided parental self-help FBT (weak recommendation). For caregivers of children and adolescents with AN, BN, or OSFED, these guidelines recommend: 1) virtual parent meal support training (weak recommendation), and 2) moderated online caregiver forums and support groups (weak recommendation). For emerging adults, these guidelines recommend: 1) CBT-based virtual group therapy for BN (strong recommendation), 2) internet CBT-based guided self-help for AN, BN, BED, and EDNOS (strong recommendation), 3) internet-based MANTRA guided self-help for AN (strong recommendation), 4) telehealth relapse prevention using MANTRA for AN (weak recommendation), and 5) guided CBT-based smartphone apps as treatment adjuncts for AN.

For caregivers of emerging adults, these guidelines recommend: 1) guided parental self-help CBT for AN or BN (strong recommendation) and 2) unguided caregiver psychoeducation self-help for most EDs (weak recommendation). Future research is required to understand the impact of sex, gender, race, socioeconomic status, and other factors on virtual care among children/adolescents and emerging adults with EDs, as well as research on more intensive services such as virtual day hospitals.
ABBREVIATIONS

AGREE II – Appraisal of guidelines for research and evaluation
AN – Anorexia Nervosa
ARFID – Avoidant/Restrictive Food Intake Disorder
BED – Binge Eating Disorder
BIAQ – Body Image Avoidance Questionnaire
BIB-GSH – Bibliotherapy guided self-help
BMI – Body mass index
BN – Bulimia Nervosa
BSQ – Body Shape Questionnaire
BULIT-R – Bulimia Test-Revised
CBT – Cognitive behavioural therapy
CBT-E – Enhanced cognitive behavioural therapy
CRT – Cognitive remediation therapy
DASS-21 – Depression, Anxiety, and Stress Scales
DEBQ-R – Dutch Eating Behaviour Questionnaire Restraint Scale
ECHO – Experienced Carers Helping Others
ED – Eating disorder
EDE-Q – Eating Disorder Examination-Questionnaire
EDI – Eating Disorders Inventory
EDNOS – Eating Disorder Not Otherwise Specified
FBT – Family-based treatment
FEAST – Families Empowered and Supporting Treatment of Eating Disorders
GDP – Guideline development panel
GRADE – Grading of Recommendations, Assessment, Development, and Evaluation system
GUIDE-M – Guideline implementability for decision excellence model
INT-GSH – Internet guided self-help
iMANTRA – internet-based Maudsley Model of Anorexia Nervosa Treatment for Adults
MANTRA – Maudsley Model of Anorexia Nervosa Treatment for Adults
MET – Motivational Enhancement Treatment
OAO – Overcoming Anorexia Online
OSFED – Other Specified Feeding and Eating Disorder
PRISMA – Preferred reporting items for systematic reviews and meta-analyses
RCT – Randomized controlled trial
SD – Standard deviation
SEED – Short Evaluation of Eating Disorders
VIA – Virtual Intervention for Anorexia Nervosa
WHO – World Health Organization

DECLARATIONS

Ethics approval and consent to participate
Consent for publication

Not applicable.

Availability of Data and Materials

Not applicable.

Competing Interests

The authors declare that they have no competing interests.

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Author contributions

JC conceived the idea for this project with input from CM, GM, LI, MN, WP, MB, MK, SF, NB, and CW. JC was primarily responsible for the overall project design, oversight of the project, and drafting of the manuscript and recommendations. NB designed and executed comprehensive searches in the databases. Several authors participated in screening abstracts and full text articles including JC, DP, and CW. DP assisted JC in the drafting of the manuscript. MB led the consensus discussions of the GDP and oversaw conflict-of-interest disclosures and management. Panel members who contributed to the drafting and finalization of the recommendations included: JC, DP, CM, SF, LI, MK, GM, MN, WS, CW,
WP, RL, TL, EW, CF, AB, CS, JG, AL, SG, MJ, GD, KB, DP, JC, AL, DK, NJ. All authors read and edited the manuscript and approved the final manuscript.

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Figure 1. PRISMA flow diagram for the impact of COVID-19 on children/adolescents and emerging adults with EDs, as well as their caregivers and clinicians.
Figure 2. PRISMA flow diagram for virtual care and eating disorders for children/adolescents, emerging adults, and their caregivers.
### Table 1. Telehealth FBT for children and adolescents

<table>
<thead>
<tr>
<th>№ of studies</th>
<th>Study design</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
<th>Indirectness</th>
<th>Imprecision</th>
<th>Other considerations</th>
<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
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<tbody>
<tr>
<td>2</td>
<td>observational studies</td>
<td>very serious&lt;sup&gt;a,b&lt;/sup&gt;</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>strong association&lt;sup&gt;c&lt;/sup&gt;</td>
<td>1 open trial (51) and 1 case report (52) with children with AN (total n=11). Significant weight gain was achieved in both studies, from baseline to end of treatment and/or at follow-up.</td>
<td>⬤⬤⬤◯ MODERATE</td>
<td>CRITICAL</td>
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<table>
<thead>
<tr>
<th>Outcomes: Weight gain</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 observational studies</td>
</tr>
</tbody>
</table>

- **a.** No control condition  
- **b.** No randomization  
- **c.** The open trial (51) had moderate to large effect sizes for participant weight gain from baseline to end of treatment and from baseline to 6-month follow-up

**Bibliography:**  
Observational studies: Open trial – Anderson 2017 (51); Case report – Goldfield 2003 (52)

### Table 2. Telehealth relapse prevention using MANTRA for emerging adults

<table>
<thead>
<tr>
<th>№ of studies</th>
<th>Study design</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
<th>Indirectness</th>
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<tbody>
<tr>
<td>1</td>
<td>open trial</td>
<td>very serious&lt;sup&gt;a,b&lt;/sup&gt;</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>none</td>
<td>1 open trial with emerging adults with AN (n=16) (53). BMI increased after the 4-month intervention.</td>
<td>⬤⬤⬤◯ LOW</td>
<td>CRITICAL</td>
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<table>
<thead>
<tr>
<th>Outcomes: BMI; EDE-Q</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 open trial</td>
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</tbody>
</table>

- **a.** No control condition  
- **b.** No randomization

**Bibliography:**
Open trial – Giel 2015 (53)

Table 3. CBT-based virtual group therapy for emerging adults

<table>
<thead>
<tr>
<th>Certainty assessment</th>
<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
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<tr>
<td># of studies</td>
<td>Study design</td>
<td>Risk of bias</td>
<td>Inconsistency</td>
</tr>
<tr>
<td>2</td>
<td>randomized trials</td>
<td>not serious</td>
<td>not serious</td>
</tr>
</tbody>
</table>

Outcomes: Body satisfaction, body attitudes, shape concerns, dietary restraint (BSQ, BIAQ, DEBQ-R)

Bibliography:
RCTs – Gollings 2006 (56), Paxton 2007 (57)

Table 4. Internet CBT-based guided self-help for emerging adults

<table>
<thead>
<tr>
<th>Certainty assessment</th>
<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
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<tbody>
<tr>
<td># of studies</td>
<td>Study design</td>
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Outcomes: ED psychopathology (SEED, EDE-Q)
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<tr>
<th>Certainty assessment</th>
<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
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<tr>
<td><strong>№ of studies</strong></td>
<td><strong>Study design</strong></td>
<td><strong>Risk of bias</strong></td>
<td><strong>Inconsistency</strong></td>
</tr>
<tr>
<td>2</td>
<td>randomized trials</td>
<td>not serious</td>
<td>not serious</td>
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</table>

Outcomes: costs (related to intervention, health care utilization, medication; assessed using Health and Labor Questionnaire)

| | | | | | | | | |
| 1 | randomized trials | not serious | not serious | not serious | none | | HIGH | CRITICAL |

Outcomes: BMI (Weight gain)
## Certainty assessment

<table>
<thead>
<tr>
<th>No of studies</th>
<th>Study design</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
<th>Indirectness</th>
<th>Imprecision</th>
<th>Other considerations</th>
<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>randomized trials</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>none</td>
<td>1 RCT with individuals with AN (n=128 VIA intervention; n=130 control) for relapse prevention (60). Intervention completers gained significantly more weight than treatment as usual controls. A 9-month follow-up of this RCT (64) (at 9-month follow-up, n=92 VIA intervention; n=120 control), very good results for BMI were seen for full completers of the intervention. Predictors for favourable course (concerning BMI) were adherence to intervention, more spontaneity, and better self-esteem.</td>
<td>HIGH</td>
<td>CRITICAL</td>
<td></td>
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</table>

### Outcomes: dropout rate

| 1             | randomized trials | not serious | not serious | not serious | none | A subsequent study (63) to the VIA RCT (60) reported VIA was well-received and highly feasible with a moderate dropout rate (15.5%). | HIGH | CRITICAL |

### Outcomes: ED symptoms (frequency of binge eating, vomiting, etc.)

| 5             | observational studies | very serious\(^a,b\) | serious\(^c\) | not serious | not serious | Strong association\(^d\) | 2 open trials (total n=228). 1 controlled study (n=31 intervention; n=31 waitlist control), 1 case series (n=38), 1 case report (n=1) all with those with BN and/or EDNOS. Both open trials had significant improvements in ED symptoms at follow-up (65, 66). In the controlled study, binge eating and vomiting abstinence rates differed significantly between the internet and control groups at post-treatment, favouring the internet group (67). The case series saw significant decreases in vomiting and weight phobia, but when bingeing and vomiting decreased, exercise increased (68). The case report did not see an improvement in ED symptoms during the intervention, although it involved ProYouth, which is used for ED prevention and early intervention (69). | LOW | CRITICAL |
### Certainty assessment

<table>
<thead>
<tr>
<th>№ of studies</th>
<th>Study design</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
<th>Indirectness</th>
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<th>Certainty</th>
<th>Importance</th>
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</thead>
<tbody>
<tr>
<td>1</td>
<td>randomized trials</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>none</td>
<td>HIGH</td>
<td>CRITICAL</td>
<td></td>
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</tbody>
</table>

1 RCT with BN and EDNOS participants (n=38 Overcoming Bulimia Online intervention; n=38 waitlist/delayed treatment control) (59). The intervention group had higher rates of cessation from binge eating and purging vs. delayed treatment condition, who experienced little change in cessation rates at follow-up. Intervention group gains were maintained or continued to improve at follow-up.

- No control condition
- No randomization
- Some discrepancies between study findings
- Large effect sizes for changes in ED symptoms scale (67) and weight phobia (68) and from baseline to post-treatment.

**Bibliography:**

RCTs – Aardoom 2016 (58), Aardoom 2017 (62), Aardoom 2016 (61), Fichter 2012 (60), Fichter 2013 (64), Fichter 2011 (63), Sanchez-Ortiz 2011 (59)

Observational studies: Open trials – Pretorius 2009 (65), Carrard 2011 (66); Controlled study – Fernandez-Aranda 2009 (67); Case series – Nevonen 2006 (68); Case report – Kindermann 2016 (69)

### Table 5. CBT-based bibliotherapy for emerging adults

<table>
<thead>
<tr>
<th>№ of studies</th>
<th>Study design</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
<th>Indirectness</th>
<th>Imprecision</th>
<th>Other considerations</th>
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<th>Importance</th>
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Outcomes: ED symptoms (frequency of binge eating, compensatory measures, etc.); ED psychopathology (EDI-2 scores)
### Table 6. Internet-based MANTRA guided self-help for emerging adults

<table>
<thead>
<tr>
<th>№ of studies</th>
<th>Study design</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
<th>Indirectness</th>
<th>Imprecision</th>
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<tbody>
<tr>
<td>3</td>
<td>randomised trials</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>none</td>
<td>1 RCT (72) and its subsequent studies (73, 74) with emerging adults with BN and EDNOS (total n=70 INT-GSH; total n=56 BIB-GSH). ED symptoms (objective binge eating and compensatory behaviour) improved significantly in both groups. Greatest improvements in ED symptoms were after 4 months; by month 18, 14.6% (7/48) of INT-GSH participants and 25% (7/28) of BIB-GSH participants were abstinent from binge eating and compensatory measures. There were no significant differences in outcomes found between the 2 groups. In both groups, lower frequency of binge eating at baseline predicted good outcomes at long-term follow-up (18 months).</td>
<td>☃️♥️♥️ HIGH</td>
<td>CRITICAL</td>
</tr>
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</table>

There were no group differences (between INT-GSH and BIB-GSH) in EDI-2 subscales (72).

Bibliography:
RCTs – Wagner 2013 (72), Wagner 2013 (73), Wagner 2015 (74)
### Certainty assessment

<table>
<thead>
<tr>
<th>№ of studies</th>
<th>Study design</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
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<th>Other considerations</th>
<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>randomised trial</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>none</td>
<td>1 RCT with AN participants (n=24 internet-MANTRA + treatment as usual; n=17 treatment as usual) (76). At 6 months: little difference between iMANTRA group (internet-based self-help workbook + email support by therapist for AN relapse prevention) and treatment as usual group. At 12 months: iMANTRA group had a higher BMI than treatment as usual group.</td>
<td>⬤⬤⬤ HIGH</td>
<td>CRITICAL</td>
</tr>
</tbody>
</table>

At 12 months, iMANTRA group had lower DASS-21 scores compared to treatment as usual group (76).

Bibliography:
RCT – Schmidt 2017 (76)

### Table 7. Online guided parental self-help – FBT for caregivers of children/adolescents

<table>
<thead>
<tr>
<th>№ of studies</th>
<th>Study design</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
<th>Indirectness</th>
<th>Imprecision</th>
<th>Other considerations</th>
<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>case series</td>
<td>very serious&lt;sup&gt;a,b&lt;/sup&gt;</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>strong association&lt;sup&gt;c&lt;/sup&gt;</td>
<td>1 case series with 19 families caring for adolescents with AN. At the end of the treatment (guided caregiver FBT training) and at follow-up, the adolescents experienced weight gain similar to clinician delivered FBT programs (large effect size) (102).</td>
<td>⬤⬤⬤ MODERATE</td>
<td>CRITICAL</td>
</tr>
</tbody>
</table>

ED-related psychopathology (EDE-Q scores) for those with AN improved by the end of the treatment (102).
Table 8. Virtual parent meal support training for caregivers of children/adolescents

<table>
<thead>
<tr>
<th>No of studies</th>
<th>Study design</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
<th>Indirectness</th>
<th>Imprecision</th>
<th>Other considerations</th>
<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>mixed methods</td>
<td>very serious&lt;sup&gt;a,b&lt;/sup&gt;</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>none</td>
<td>1 open trial with adolescents with, at risk, or at high-risk for AN (n=12 diagnosed with AN, n=12 at risk for AN, n=22 at high risk for AN) (103). At the end of the 6 family-based early intervention online sessions, adolescents remained stable or increased in ideal body weight (weight gain).</td>
<td>☒ ☒ ☐ ☐ LOW</td>
<td>CRITICAL</td>
</tr>
</tbody>
</table>

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<table>
<thead>
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<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>a.</td>
<td>No control condition</td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>b.</td>
<td>No randomization</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>c.</td>
<td>Large effect size from baseline to end of treatment and baseline to follow-up (102)</td>
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<td></td>
</tr>
</tbody>
</table>

Bibliography:
Case series – Lock 2017 (102)
Open trial – Jones 2012 (103)
# Table 9: Unguided caregiver self-help using web-based and/or pre-recorded videos focused on psychoeducation and communication skills for caregivers of emerging adults

<table>
<thead>
<tr>
<th>Certainty assessment</th>
<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
</tr>
</thead>
<tbody>
<tr>
<td>№ of studies</td>
<td>Study design</td>
<td>Risk of bias</td>
<td>Inconsistency</td>
</tr>
<tr>
<td>1</td>
<td>randomized trial</td>
<td>not serious</td>
<td>not serious</td>
</tr>
<tr>
<td>1</td>
<td>randomized trial</td>
<td>not serious</td>
<td>not serious</td>
</tr>
</tbody>
</table>

Outcomes: accommodation and enabling scale for EDs; caregiver burden

Bibliography:
RCTs – Dimitropoulos 2019 (105), Quadflieg 2017 (106)
### Table 10. Guided caregiver self-help – Skills for caregivers of children and adolescents

<table>
<thead>
<tr>
<th>№ of studies</th>
<th>Study design</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
<th>Indirectness</th>
<th>Imprecision</th>
<th>Other considerations</th>
<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>mixed methods</td>
<td>very serious&lt;sup&gt;a&lt;/sup&gt;</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>none</td>
<td><img src="https://placekitten.com/150/150" alt="⊕⊕⨁⨁" /> LOW CRITICAL</td>
<td>1 mixed methods study with caregivers (n=16) for children with AN and BN (107). Caregivers expressed high levels of satisfaction with most aspects of the DVD and coaching skills training. From baseline to post-intervention, caregiver general distress (General Health Questionnaire scores) decreased significantly.</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> No control condition  
<sup>b</sup> No randomization

**Bibliography:**  
Mixed methods – Sepulveda 2008 (107)

### Table 11. Guided parental self-help CBT (skills training approach + workbook or CBT-based online modules) for caregivers of emerging adults

<table>
<thead>
<tr>
<th>№ of studies</th>
<th>Study design</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
<th>Indirectness</th>
<th>Imprecision</th>
<th>Other considerations</th>
<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>randomised trials</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>none</td>
<td><img src="https://placekitten.com/150/150" alt="⊕⊕⊕⊕" /> HIGH CRITICAL</td>
<td>1 RCT with caregivers (n=134 in telephone coaching ECHO intervention; n=134 control) for individuals with AN (76). Caregivers in the intervention group had reduced expressed emotion levels at patient discharge and 6-month follow-up.</td>
<td></td>
</tr>
</tbody>
</table>

Caregivers in the telephone coaching intervention group experienced greater reductions in ED symptom impact scale scores than in the control group (76).
<table>
<thead>
<tr>
<th>№ of studies</th>
<th>Study design</th>
<th>Risk of bias</th>
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<th>Indirectness</th>
<th>Imprecision</th>
<th>Other considerations</th>
<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
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<td>randomised trials</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>none</td>
<td>1 RCT with caregivers for individuals with AN (n=33 in OAO web-based intervention; n=30 in telephone and email hotline control) (108). Compared with control, the OAO intervention significantly reduced anxiety and depression in caregivers at post-treatment; caregivers in the OAO intervention had greater reductions in expressed emotion than controls, but these were not significant.</td>
<td>⬤⬤⬤⬤ HIGH</td>
<td>CRITICAL</td>
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<td>Similar to expressed emotion, caregivers’ ED symptom impact scale scores were also reduced across both groups, but there was no significant difference between groups (108).</td>
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<tr>
<td>1</td>
<td>randomised trials</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>none</td>
<td>1 RCT with caregivers for individuals with AN (n=19 OAO + guidance intervention; n=18 OAO only) (109). Levels of expressed emotion reported by caregivers at post-intervention were reduced but did not differ significantly between the groups. Those with AN did not perceive that their caregivers’ levels of expressed emotion had significantly changed.</td>
<td>⬤⬤⬤⬤ HIGH</td>
<td>CRITICAL</td>
</tr>
<tr>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>Caregivers’ ED symptom impact scale scores were also reduced across both groups, but there was no significant difference between groups (109).</td>
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</tbody>
</table>

Bibliography:
RCTs – Schmidt 2017 (76), Grover 2011 (108), Hoyle 2013 (109)
Table 12. Moderated online caregiver forums and teleconferenced caregiver support groups for caregivers of children/adolescents and emerging adults

<table>
<thead>
<tr>
<th># of studies</th>
<th>Study design</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
<th>Indirectness</th>
<th>Imprecision</th>
<th>Other considerations</th>
<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>open trial</td>
<td>very serious(^{a,b})</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>none</td>
<td>1 open trial with 13 caregivers with a child with AN (111). The virtual chat room sessions led by an FBT-trained therapist had high satisfaction ratings (91.7%; feasibility and acceptability) among caregivers. Caregivers reported looking forward to the chat sessions and that they were accessible, convenient, and easy to use.</td>
<td>⬤⤤⤤⤤</td>
<td>LOW</td>
</tr>
<tr>
<td>1</td>
<td>case report</td>
<td>very serious(^{a,b})</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>none</td>
<td>1 case report with 6 caregivers with a child with an ED, participating in teleconferencing group sessions to foster caregiver support (112). Monthly in-person meetings were added as more participants became interested. Overall, caregivers in the teleconferencing support groups felt they were helpful for those who feel emotionally isolated (feasibility and acceptability), but in-person meetings for support may be more ideal.</td>
<td>⬤⤤⤤⤤</td>
<td>LOW</td>
</tr>
</tbody>
</table>

\(^{a}\) No control condition  
\(^{b}\) No randomization

Bibliography:  
Open trial – Binford Hopf 2013 (111)  
Case report – Marx 2006 (112)
### Table 13. Smartphone apps for emerging adults

<table>
<thead>
<tr>
<th>Certainty assessment</th>
<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
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<td>randomized trials</td>
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<td></td>
</tr>
<tr>
<td>Outcomes: ED psychopathology (EDE-Q); BMI</td>
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<td></td>
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<tr>
<td>Outcomes: ED behaviours (restrictive eating, binge eating, compensatory measures)</td>
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</tbody>
</table>

a. No control condition  

b. No randomization
Table 14. In person care versus home monitoring for children and adolescents

<table>
<thead>
<tr>
<th>№ of studies</th>
<th>Study design</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
<th>Indirectness</th>
<th>Imprecision</th>
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<th>Impact</th>
<th>Certainty</th>
<th>Importance</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>open trial</td>
<td>very serious&lt;sup&gt;a,b&lt;/sup&gt;</td>
<td>not serious</td>
<td>not serious</td>
<td>not serious</td>
<td>none</td>
<td></td>
<td>LOW</td>
<td>CRITICAL</td>
</tr>
</tbody>
</table>

**Outcomes: Vital signs (heart rate and heart rate variability)**

1 open trial comparing adolescent girls with AN using wearable sensors to monitor heart rate and heart rate variability (n=27) versus healthy controls (n=15) (113). Results showed that the wearable sensors were feasible to monitor vital signs in adolescents with AN, but by a clinician rather than a caregiver.

**Outcomes: Monitoring weight at home**

1 case report in a descriptive study with 1 child with AN (37). The family utilized telehealth for FBT during COVID-19 where the child’s caregivers successfully measured the child’s weight at home using a home scale, as well as blood pressure and heart rate and reported these measures to the clinician.

---

a. No control condition
b. No randomization

Bibliography:
Open trial – Billeci 2015 (113)
Case report – Wood 2020 (37)