Neurokinin 3 Receptor Antagonists Compared With Serotonin Norepinephrine Reuptake Inhibitors for Non-Hormonal Treatment of Menopausal Hot Flushes: A Systematic Qualitative Review

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ABSTRACT

Introduction: Hot flushes/flashes (HFs) or other vasomotor symptoms affect between 45 and 97% of women during menopause. Hormone replacement therapy (HRT) is effective at alleviating menopausal symptoms, but some women cannot or prefer not to take HRT. Since current non-hormonal options have suboptimal efficacy/tolerability, there is a pressing need for an effective, well-tolerated alternative. The neurokinin 3 receptor (NK3R) has recently been implicated in the generation of menopausal HFs and represents a novel therapeutic target to ameliorate HF symptoms. This review aims to assess if NK3R antagonists (NK3Ras) are more effective than Serotonin Norepinephrine Reuptake Inhibitors (SNRIs)—currently a common choice for non-hormonal treatment of menopausal HFs.

Methods: Studies were identified after systematically searching Ovid MEDLINE and EMBASE databases based on PRISMA guidelines. Trial quality and bias were assessed. Key efficacy outcomes (HF frequency, HF severity and number of night-time awakenings/night-sweats) and selected safety outcomes were extracted and analysed.

Results: Seven SNRI and four NK3Ra placebo-controlled randomised trials (plus four follow-up reports) were included in this review. NK3Ra administration resulted in a larger reduction from baseline in HF frequency, HF severity and night-sweats compared to SNRIs. Five of seven SNRI trials showed a reduction in HF frequency that was statistically significant (by 48–67% from baseline at weeks 8 or 12) whereas all NK3Ra trials showed a statistically significant reduction in HF frequency (by 62–93% from baseline at weeks 2, 4 or 12). While SNRI trials reported poor tolerability, particularly nausea, NK3Ra trials reported good tolerability overall, although two trials reported elevation in transaminases.

Conclusion: NK3Ras trials show encouraging efficacy and tolerability/safety. Completion of phase 3 NK3Ra trials are required to confirm efficacy and uphold safety/tolerability data but
phase 2 results suggest that NK3Ras are more effective than SNRIs for non-hormonal treatment of menopausal HFs.

**Keywords:** Menopause; Vasomotor symptoms; Hot flushes/flashes; Neurokinin 3 receptor antagonist; Fezolinetant; Elinzanetant (NT-814); MLE4901; Serotonin Norepinephrine Reuptake Inhibitor; Venlafaxine; Desvenlafaxine

### Key Summary Points

Between 45 and 97% of menopausal women suffer vasomotor symptoms (hot flushes/flashes and night-sweats) which can significantly impact their quality of life and 10–20% find them almost intolerable.

Conventional treatment is hormone replacement therapy (HRT) but some women cannot or prefer not to take HRT. However, current non-hormonal options have suboptimal efficacy and tolerability.

Neurokinin B, predominantly acting via the neurokinin 3 receptor (NK3R), has been implicated in the generation of menopausal hot flushes/flashes.

We undertook a systematic qualitative review to compare outcomes of placebo-controlled randomised clinical trials using neurokinin 3 receptor antagonists (NK3Ras) with those using Serotonin Norepinephrine Reuptake Inhibitors (SNRIs) for the non-hormonal treatment of menopausal hot flushes/flashes.

NK3Ra trials reported numerically superior efficacy and better safety/tolerability compared with SNRIs trials. Provided that phase 3 NK3Ra trials are supportive, NK3Ras appear a promising therapy for this challenging area.

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### INTRODUCTION

Hot flushes/flashes (HFs), defined as transient sensations of heat, flushing, sweating and chills, affect between 45 and 97% of women during menopause and can significantly degrade their quality of life [1]. HFs often lead to sleep disturbances for a prolonged period, with recurrent episodes occurring over a median duration of 7.4 years [2]. Between 10 and 20% of women find HFs almost intolerable [3]. While the full pathophysiological mechanism of HFs remains elusive, oestrogen deficiency appears to play a causative role. Hormone replacement therapy (HRT) remains the most commonly prescribed treatment to alleviate menopausal symptoms provided there are no contraindications (Table 1) [4]. However, the use of HRT has decreased after reports of increased risk of cardiovascular (CV) disease, breast cancer, stroke and pulmonary embolism [5]. While subsequent data suggest that the benefits of HRT typically outweigh the risks for women without contraindications, many now seek non-hormonal alternatives.

**Table 1** Typical contraindications for hormone replacement therapy [4]

| Contraindications for hormone replacement therapy |
|---------------------------------------------------|
| Current, past, or suspected breast cancer          |
| Known or suspected oestrogen-dependent cancer      |
| Undiagnosed vaginal bleeding                       |
| Untreated endometrial hyperplasia                  |
| Previous idiopathic or current venous thromboembolism (deep vein thrombosis or pulmonary embolism), unless the woman is already on anticoagulant treatment |
| Active or recent arterial thromboembolic disease (for example, angina or myocardial infarction) |
| Active liver disease with abnormal liver function tests |
| Pregnancy                                          |
| Thrombophilic disorder                             |
Current non-hormonal treatment options include Serotonin Norepinephrine Reuptake Inhibitors (SNRIs) or Selective Serotonin Reuptake Inhibitors (SSRIs), clonidine, gabapentin and pregabalin. Of these, SNRIs/SSRIs are probably the most commonly recommended

Fig. 1 Relationship between KNDy neurons, GnRH neurons, and the heat-defence pathway. KNDy neurones in the infundibular nucleus secrete the neuropeptides Kiss (encoded by the KISS1 gene), NKB (encoded by the TAC3 gene) and Dyn (encoded by the PDYN gene). NKB and Dyn act autosynaptically, stimulating and inhibiting further Kiss release, respectively. Kiss acts on GnRH neurones. GnRH stimulates LH and FSH release from the anterior pituitary which stimulates ovarian sex steroid production. During menopause, lack of oestadiol (E2) negative feedback results in increased expression of KISS1 and TAC3 mRNA but decreased expression of PDYN mRNA. Consequently, KNDy neurones become hypertrophied, as seen by increased size of nuclei/nucleoli and increased Nissl substance. KNDy neurones project to the hypothalamic thermoregulatory centre (the median POA and adjacent MnPO). During menopause, the increase in NKB signalling and overstimulation of KNDy neurones increases activity in the thermoregulatory centre which then becomes hypersensitive to external cues from peripheral sensors, leading to activation of heat dissipation effectors. KNDy kisspeptin–neurokinin B–dynorphin, Kiss kisspeptin, NKB neurokinin B, Dyn dynorphin, GnRH gonadotropin-releasing hormone, LH luteinizing hormone, FSH follicle-stimulating hormone, POA pre-optic area, MnPO median preoptic nucleus, E2 oestradiol, ERα oestrogen receptor alpha, PR progesterone receptor, KISS1 kisspeptin gene, TAC3 tachykinin 3 gene, PDYN prodynorphin gene. The figure was created with BioRender.com
[6], with data for venlafaxine considered the most convincing [7]. Nevertheless, none of the current non-hormonal drugs have optimal efficacy and use in clinical practice may be limited by side effects or by interactions with other medications (for example, paroxetine or other CYP2D6 inhibitors should not be co-prescribed with tamoxifen [7] which is clinically relevant considering that a key indication for a non-hormonal treatment is prior breast cancer). Therefore, there remains a pressing need for an effective and well-tolerated non-hormonal alternative.

The hypothalamic neuropeptide kisspeptin (Kiss; encoded by the KISS1 gene) is required for human fertility being a potent stimulator of the hypothalamic–pituitary–gonadal axis (Fig. 1) [8]. In primates, a population of Kiss neurones located in the hypothalamic infundibular nucleus co-express the neuropeptides, neurokinin B (NKB) and dynorphin (Dyn), and are thus termed kisspeptin–neurokinin B–dynorphin (KNDy) neurones [9]. NKB [encoded by the tachykinin 3 (TAC3) gene] and Dyn [encoded by the prodynorphin (PDYN) gene] regulate pulsatile Kiss secretion from KNDy neurones by acting auto-synaptically via neurokinin 3 receptors (NK3R) and Dyn (kappa opioid) receptors to stimulate and inhibit the release of Kiss, respectively [10]. KNDy neurones project to the hypothalamic thermoregulatory centre in mammals (the median preoptic area and adjacent median preoptic nucleus) [11]. Gonadal feedback is transmitted by oestrogen receptor alpha (ERα) and progesterone receptors (PR) expressed on Kiss neurones. KNDy neurones are normally stimulated by NK3R activation and inhibited by oestrogen. However, during menopause declining oestrogen levels leads to NKB hypersecretion, and overstimulation of KNDy neurones in particular. This elicits increased activity in the thermoregulatory centre, which shifts the thermoneutral zone increasing sensitivity to external cues and triggering more frequent heat dissipation responses such as HFs [12].

Blockade of NKB signalling with the use of an NK3R antagonist (NK3Ra) is proposed to normalise KNDy neurone activity and thus may help alleviate HFs in menopausal women. This review aimed to (1) evaluate randomised controlled trials (RCTs) reporting the efficacy and tolerability of NK3RAs for the treatment of menopausal HFs and (2) to put these findings in clinical context by qualitative comparison with outcome data from RCTs using the SNRI venlafaxine (or the succinate salt of its active metabolite, desvenlafaxine) to treat menopausal HFs.

**METHODS**

To put the clinical outcomes of the NK3Ra studies into context, a clinically relevant alternative non-hormonal therapy was required for comparison. While arguments could be made for selecting any one of the SNRIs/SSRIs, clonidine, gabapentin or pregabalin, multiple guidelines and reviews suggest that SSRIs/SNRIs are the most commonly recommended [6], with trial evidence strongest for venlafaxine/desvenlafaxine, paroxetine and citalopram/escitalopram [13]. Of these, venlafaxine (or its active metabolite desvenlafaxine) was selected for 3 reasons: modest current use was confirmed after consultation with each of 4 UK regional menopause clinics (London, Edinburgh, Cardiff and Belfast), a College report considered the most convincing SSRI/SNRI efficacy data were for venlafaxine [7], and, since venlafaxine does not interact with tamoxifen, it is the preferred treatment for breast cancer survivors taking tamoxifen [14], a key indication for a non-hormonal menopause treatment. In contrast, the SSRI paroxetine is a potent CYP2D6 inhibitor and is contra-indicated with tamoxifen. The SSRI fluoxetine and SNRI duloxetine are also CYP2D6 inhibitors (albeit more moderate).

A systematic search of the published literature up to 17/2/2021 was undertaken on Ovid MEDLINE and EMBASE, based on PRISMA guidelines [15] and using the following key words: “venlafaxine”, “desvenlafaxine”, “hot flushes”, “hot flashes”, “vasomotor symptoms”, “VMS”, “menopause”, “neurokinin B”, “dynorphin”, “kisspeptin”, “kappa”, “neurokinin 1”, “neurokinin 3”, “neurokinin B”. Further studies were identified by cross-referencing references from the qualifying studies. Duplicates were
removed and studies were screened using inclusion/exclusion criteria.

Inclusion criteria were: studies of peri-menopausal, menopausal and postmenopausal women (natural or surgically induced only), having HFs, receiving NK3Ra or venlafaxine/desvenlafaxine treatment, comparison with a placebo arm, frequency and severity of HFs reported, primary research design and in English language.

Exclusion criteria were: pre-menopausal women, chemically or medically induced menopause, not having HFs, not receiving treatment with NK3Ra or venlafaxine/desvenlafaxine, or receiving treatment for HFs in addition to NK3Ra or venlafaxine/desvenlafaxine. Qualifying studies were required to meet all inclusion criteria and studies discarded if they met any one of the exclusion criteria.

Only women with natural or surgically-induced menopause were included to ensure a definitive and irreversible diagnosis. Women with medically-induced menopause (such as chemotherapy or use of GnRH analogues) were excluded, as their ovarian function may have had the potential to recover (and reverse menopause). Men were excluded due to the different physiology involved in the development of HFs.

Women concurrently receiving any other medications to relieve HFs other than the study drugs were excluded so that the measured effect could be attributed solely to the drugs under investigation.

A large placebo effect is well described in HF treatment trials [16], thus only studies comparing the active drug versus placebo were selected so that the drug's effect could be evaluated in context.

Once qualifying studies were identified, study quality was assessed using Critical Appraisal Skills Programme (CASP) methodology [17]. The CASP method evaluated four main components (study design, methodology, reporting of results and clinical implications), with each component addressed by either 2 or 3 questions. The Cochrane Collaboration's tool [18] was used to assess for risk of bias in each study, comprising 7 questions defined by the Cochrane handbook to evaluate selection (randomisation or allocation concealment), reporting, performance, detection, attrition and other bias. One author (S.J.M.) undertook the primary detailed evaluation of quality and bias for each study. Equivocal findings were discussed with a second author (J.A.T.) to reach consensus.

To enable comparison of SNRI and NK3Ra outcomes, while studies may include different outcome measures [19], each qualifying study was assessed for the specific key efficacy outcomes of HF frequency, HF severity and, if stated, night-time awakenings due to HFs, plus safety/tolerability outcomes including common adverse events (AEs), serious adverse events (SAE) and discontinuation due to AEs. Since SNRI studies were phase 3 whereas NK3Ra trials were phase 2, a systematic qualitative review rather than a meta-analysis was considered the most appropriate.

This paper is based on previously published data and does not involve any new studies with human or animal subjects performed by any of the authors.

RESULTS

Figure 2 illustrates the PRISMA flow diagram summarising the search process. The initial search yielded 651 records. After 127 duplicates were removed, 524 studies were screened for inclusion (based on the inclusion/exclusion criteria). A total of 500 studies were excluded based on the title or abstract. The remaining 24 studies required assessment of the full text, and a further three studies were found by cross-referencing references from these studies. This resulted in 15 qualifying studies for qualitative synthesis (seven SNRI trials plus two follow-up papers [20–28] and four NK3Ra trials plus two follow-up papers [29–34]).

Table 2 summarises the CASP quality assessment for each study. For illustrative purposes, when assessing the four components, if all answers were ‘Yes’, the score was designated as ‘high’ quality of evidence (green). If one or more answers was ‘No’, the score was assigned as ‘low’ quality of evidence (red). If one or more answers was ‘cannot tell’ (but none was ‘no’)

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the score was assigned as ‘unclear’ quality of evidence (orange). The full quality analysis for each study is available upon request. Overall, the studies were mostly of high quality.

Table 3 summarises the Cochrane Collaboration’s tool bias assessment for each study. For illustrative purposes, a low risk of bias subtype was shown as green, a high risk as red and an unclear risk as orange. The full quality analysis for each study is available upon request. Overall, the studies were mostly of low bias.

Table 4 summarises the study design, efficacy and tolerability/safety outcomes for SNRI trials. Overall, the SNRI trials showed modest HF reduction. There was a statistically significant
reduction in HF frequency versus placebo in five of seven trials [21–23, 25, 27] (48–67% reduction from baseline at week 8 or 12 in the setting of a 25–51% placebo response) and a statistically significant reduction in HF severity versus placebo in five of seven trials [21–23, 25, 27] for at least one dose and one time point (24–31% reduction from baseline at week 12 in the setting of a 12–18% placebo response where percentage data were reported). Night-time awakenings due to HFs showed a statistically significant reduction in the three trials [21–23] that measured this outcome (53–77% reduction from baseline at week 12 in the setting of 44–63% placebo response). However, HF frequency and HF severity did not show a statistically significant reduction versus placebo in two SNRI trials [20, 24]. SNRI-treated subjects experienced more AEs (in all trials) and discontinuations due to AEs (in all but one trial [22]) versus placebo. Common AEs included nausea, dry mouth, insomnia, dizziness and constipation (with nausea being reported to be the most debilitating).

Table 5 summarises the study design efficacy and tolerability/safety outcomes for NK3Ra trials.
Overall, in comparison to SNRI trials, the NK3Ra trials reported a larger reduction versus baseline in HF outcomes. NK3Ras led to a statistically significant reduction in HF frequency versus placebo at all time points, across all four trials, apart from the two lowest doses in the dose-ranging study by Trower et al. [34] (62–93% reduction from baseline at weeks 2, 4 or 12 in the setting of a 28–55% placebo response). HF severity also showed a reduction versus placebo in all trials (41–94% reduction from baseline measured at weeks 2, 4 or 12 in the setting of a 5–46% placebo response), although not all doses reached significance at all time points in the two dose-ranging trials [32, 34]. The number of night-time HF/frequency of waking due to night-sweats showed a statistically significant reduction versus placebo in the two trials [30, 34] that measured this outcome (63–81% reduction from baseline at weeks 2 and 4 in the setting of a 22–32% placebo response).

**DISCUSSION**

In this qualitative analysis, NK3Ras reported larger reductions from baseline compared to SNRIs in terms of HF frequency, HF severity and night-sweats. All four NK3Ra trials reported statistically significant reductions in HF frequency by 62–93% from baseline at weeks 2, 4 or 12 (in the setting of a 28–55% placebo response), whereas only five of seven SNRI trials reported reduced HF frequency by 48–67% from baseline at weeks 8 or 12 (in the setting of a 25–51% placebo response) and two of seven trials reported no significant reduction. NK3Ra trials reported generally good tolerability (although transaminases elevation was noted in two trials) whereas SNRI trials reported less favourable tolerability, with nausea being common.

**Trial Strengths and Weaknesses**

While both the SNRI and NK3Ra trials had many strengths, potential limitations were
| Clinical trial | Trial design and key participant BL data | Key outcomes | Results (p values vs. placebo) |
|----------------|------------------------------------------|--------------|--------------------------------|
| 1 Evans et al  | 12-week RCT: 1 US centre                 | 1\%:         | 1\%: absolute or \% data not reported |
| Obstetrics and | Mean age = 52.15 years                   | # of mild, mod, severe, and very severe HFs/day | No sig. ↓ in HF # vs. placebo (p = 0.2) |
| Gynecology     | 80 randomised (1:1) to venlafaxine XR (37.5 mg/day for 1 week, titrated to 75 mg/day for 11 weeks) vs. placebo | Avg. HF severity | No sig. ↓ in mean HF severity (p value not given) |
| (2005) [20]    |                                           | 2\%:         | 2\%:  
|                |                                           | Important AEs/SAEs | AEs ↑ in dry mouth (81% vs. 44%), ↑ in sleeplessness (88% vs. 47%) and ↓ appetite (81% vs. 53%) vs. placebo |
|                |                                           | Discontinuation due to AEs | ↑ withdrawals [11 vs. 8 (p = ns)] due to difficulty sleeping, ↓ libido, nausea or anxiety |
|                |                                           | 1\%:         | 1\%:  
|                |                                           | from BL in avg. daily # of mod/severe HFs at week 4 and 12 | ↓ from BL in avg. daily # of mod/severe HFs vs. placebo (No ↓ with 50 mg or 200 mg) |
|                |                                           | from BL in avg. daily HF severity score at week 4 and 12 | 100 mg ↓ at week 4 (−6.62 vs. −5.22; p = 0.013) and week 12 (−7.23 [64%]) vs. −5.50 (51%); p = 0.005 |
|                |                                           | from BL in daily # of night-time awakenings due to HFs at week 4 and 12 | 150 mg ↓ at week 12 (−6.94 [60%]) vs. −5.50 (51%); p = 0.020, but not at week 4 |
|                |                                           | Important AEs/SAEs | 2\%:  
|                |                                           | Discontinuation due to AEs | ↓ from BL in avg. daily HF severity scores vs. placebo (No ↓ with 50 mg or 150 mg) |
|                |                                           | 150 mg and 200 mg: AEs during week 1 only (both p < 0.05), most commonly, nausea, dry mouth, insomnia | 100 mg ↓ at week 12 (−0.80 [31%]) vs. −0.47 [18%]; p = 0.002, but not at week 4 |
|                |                                           | 200 mg ↓ at week 12 (−0.74 [27%]) vs. −0.47 [18%]; p = 0.013, but not at week 4 | 2\%:  
|                |                                           | Daily # of night-time awakenings vs. placebo at week 12 (week 4 not reported) (No ↓ with 50 mg) | 100 mg, 150 mg and 200 mg ↓ at week 12 (−2.77/night [76.9%]; p = 0.013), −2.69/night [69.9%]; p = 0.034 and −2.68/night [70.5%]; p = 0.043, respectively vs. −2.21/night [63.4%]) |
|                |                                           | AEs vs. placebo | AEs possibly Tx-related: 2 increased LFTs, 1 cholecystitis |
|                |                                           | 150 mg and 200 mg ↑ AEs during week 1 only (both p < 0.05), most commonly, nausea, dry mouth, insomnia (No ↑ with 50 mg and 100 mg) | ↑ hypertension (5.9% overall vs. 1.3% placebo; p = ns), 5 vs. 0 CV events at 1 year (p = ns) |
|                |                                           | ↑ discontinuations due to AEs 150 and 200 mg during week 1 only (both p < 0.001). (No ↑ with 50 mg and 100 mg) | 3 SAEs possibly Tx-related: 2 increased LFTs, 1 cholecystitis |

*Table 4* Serotonin-norepinephrine reuptake inhibitor (SNRI) study results
### Table 4 continued

| Clinical trial | Trial design and key participant BL data | Key outcomes | Results (p values vs. placebo) |
|----------------|------------------------------------------|--------------|--------------------------------|
| 3              | 12 week DBRCT; 34 US centres            |              |                                |
| Archer et al   | Mean age = 53.36 years (29–71)          |              |                                |
| American Journal of Obstetrics and Gynecology (2009) [22] | Mean BMI = 27.86 kg/m² (17.2–40.1) |              |                                |
|                | 458 randomised to desvenlafaxine 100 mg/day or 150 mg/day vs. placebo for 12 weeks [50 mg/day for 3 days, titrated to 100 mg/day on day 4 (titrated to 150 mg/day on day 8 for 150 mg/day group)] |              |                                |
|                | 2 week dose-tapering                      |              |                                |
|                | 436 included in mITT analysis            |              |                                |
|                |                                          | 1\*:         |                                |
|                | Δ from BL in avg. daily # of mod/severe HFs at weeks 4 and 12 |              |                                |
|                | Δ from BL in avg. daily HF severity score at weeks 4 and 12 |              |                                |
|                | Δ from BL in # of night-time awakenings at weeks 4 and 12 |              |                                |
| 1\*:           | Discontinuation due to AEs               |              |                                |
| 2\*:           | Important AEs/SAEs                       |              |                                |
|                | Discontinuation due to AEs               |              |                                |
| 1\*:           |                                          | 1\*:         |                                |
|                | daily # of HF from BL vs. placebo with 100 mg and 150 mg at week 4 (both p < 0.012, no % given) and week 12 (65.4%; p = 0.005; and 66.6%; p = 0.012, respectively vs. 50.8%) |              |                                |
| 2\*:           | daily HF severity score from BL vs. placebo with 100 mg and 150 mg at week 12 (–0.65 [27%]; and –0.66 [27.5%], respectively vs. –0.33 [13.75%]; both p < 0.001), and at all other time points |              |                                |
|                |                                          | 2\*:         |                                |
|                | daily # of night-time awakenings from BL vs. placebo with 100 mg and 150 mg at week 4 (–1.8 and –1.6, respectively vs. –1.2) and week 12 (–2.0 [60.6%] and –1.8 [58.1%], respectively vs. –1.4 [43.8%]); all p < 0.048 |              |                                |
|                |                                          |              |                                |
| 4              | 26 week DBRCT; 32 US centres            |              |                                |
| Archer et al   | Mean age = 53.7 years                    |              |                                |
| American Journal of Obstetrics and Gynecology (2009) [23] | Mean BMI = 27.1 kg/m² (15.9–40.4) |              |                                |
|                | 567 randomised to desvenlafaxine 100 mg/day or 150 mg/day vs. placebo for 26 weeks |              |                                |
|                | 484 included in mITT analysis            |              |                                |
|                |                                          | 1\*:         |                                |
|                | Δ from BL in avg. daily # of mod/severe HFs at weeks 4 and 12 |              |                                |
|                | Δ from BL in avg. daily HF severity score at weeks 4 and 12 |              |                                |
|                | Δ from BL in # of night-time awakenings due to HFs |              |                                |
| 1\*:           | Important AEs/SAEs                       |              |                                |
| 2\*:           | Discontinuation due to AEs               |              |                                |
| 1\*:           |                                          | 1\*:         |                                |
|                | daily # of HFs from BL vs. placebo at week 4 and 12 |              |                                |
|                | (week 12: 100 mg 60%\*, p = 0.002; 150 mg 66.6%\*, p < 0.001, vs. 47%\*) |              |                                |
| 2\*:           | daily HF severity score from BL vs. placebo at week 4 and 12 |              |                                |
|                | (week 12: 100 mg 69%\*, p = 0.001), whereas 100 mg did not (61%\*, p = 0.061) but study not powered for efficacy > 12 weeks |              |                                |
|                |                                          | 2\*:         |                                |
|                | daily # of night-time awakenings from BL vs. placebo at week 4 and 12 |              |                                |
|                | (week 12: 100 mg 24%\*, p = 0.002; 150 mg 29%\*, p < 0.001 vs. 13%\*), 150 mg maintained Δ at week 26 (p = 0.008, no % given) |              |                                |
|                |                                          | 2\*:         |                                |
|                | daily # of night-time awakenings from BL vs. placebo at week 4 (actual Δ not given) and week 12 [100 mg – 2.0 (52.6%\*); 150 mg – 2.4 (68.6%\*) vs. – 1.6 (47.1%\*)]; all p ≤ 0.026 |              |                                |
|                |                                          |              |                                |
|                | AE during week 1 only vs. placebo (p < 0.05), most commonly nausea (44.6%), dizziness, insomnia, dry mouth, constipation |              |                                |
|                | SAE possibly Tx-related: hypotension (1 subject with 150 mg) |              |                                |
|                | Discontinuation due to AEs no diff. [but numerically ↑ in 150 mg group (p = ns)] |              |                                |
|                | Discontinuation due to AEs               |              |                                |
Clinical trial | Trial design and key participant BL data | Key outcomes | Results (p values vs. placebo)
--- | --- | --- | ---
5 | 12-week DBRCT: 35 European centres, 2 centres in South Africa, 1 centre in Mexico | 1<sup>st</sup>: Δ from BL in avg. daily # of mod/severe HFs at weeks 4 and 12 | 1<sup>st</sup>: No ↓ in daily # of HFs from BL vs. placebo at week 4 (Δ = 4.63 vs. − 4.38, p = 0.558) and week 12 (Δ = 5.78 (37.7%) vs. − 5.82 (37.5%), p = 0.921)
Bouchard et al Climaeteric (2012) [24] | Mean age = 53.6 years (40–66 years) Mean BMI = 26 kg/m² (16–34) ≥ 485 randomised (1:1:1) to desvenlafaxine 100 mg/day, tibolone 2.5 mg/day, vs. placebo for 12 weeks 451 included in mITT analysis | 2<sup>nd</sup>: Δ from BL in avg. daily HF severity at weeks 4 and 12 2<sup>nd</sup>: Important AEs/SAEs Discontinuation due to AEs | No ↓ in daily HF severity from BL vs. placebo at week 4 (Δ = 0.37 vs. − 0.31, p = 0.352) and week 12 (Δ = 0.61 (26.8%) vs. − 0.61 (26.5%), p = 0.943)
6a | 52-week DBRCT: 122 US and Canadian centres | 1<sup>st</sup>: Δ from BL in avg. daily # of mod/severe HFs at weeks 4 and 12 | 1<sup>st</sup>: No ↓ in daily # of HFs vs. placebo at week 4 (Δ = − 6.5 HFs (55.5%) vs. − 3.6 (31%); p < 0.001) and week 12 (Δ = − 7.3 HFs (62.2%) vs. − 4.5 (38.6%); p < 0.001)
Pinkerton et al Menopause (2013) [25] | Mean age = 54 years (45–71) Mean BMI = 26.45 kg/m² (16.9–35.3) 396 randomised (1:1:1) to desvenlafaxine 100 mg/day vs. placebo for 52 weeks (50 mg/day for 1 week, titrated to 100 mg/day for 51 weeks) 2 week dose-tapering 365 included in mITT analysis Pinkerton et al. (2013) [25] reports 12-week data from an efficacy substudy (part of a larger n = 2186 safety study) | 2<sup>nd</sup>: Δ from BL in avg. daily HF severity scores at weeks 4 and 12 2<sup>nd</sup>: Important AEs/SAEs Discontinuation due to AEs | No ↓ in daily HF severity vs. placebo at week 4 (Δ = − 0.47 (20%) vs. − 0.19 (8%); p < 0.001) and week 12 (Δ = − 0.59 (25%) vs. − 0.28 (12%); p < 0.001)
6b | Pinkerton et al. (2015) [26] reports 52-week data from the same efficacy substudy population | 1<sup>st</sup>: Δ from BL in avg. daily # of HFs at weeks 12, 26, 52 | 1<sup>st</sup>: No ↓ in daily # of HFs at 12 weeks (Δ = − 7.5 HFs (64.4%) vs. − 5.0 (43.3%); p < 0.001), 26 weeks (Δ = − 8.6 HFs (74.9%) vs. − 6.3 (54%); p < 0.001 and 52 weeks (Δ = − 7.7 HFs (66%) vs. − 4.8 (41%); p < 0.001)
Pinkerton et al. Menopause (2013) [26] | See above | 2<sup>nd</sup>: Δ from BL in avg. daily HF severity scores at weeks 12, 26, 52 2<sup>nd</sup>: Important AEs/SAEs Discontinuation due to AEs | No ↓ in daily HF severity score at 12 weeks (Δ = − 0.63 (27%) vs. − 0.3 (13%); p < 0.001), 26 weeks (Δ = − 0.85 (36%); vs. − 0.53 (22%); p = 0.001) and 52 weeks (Δ = − 0.75 (32%); vs. − 0.44 (19%); p = 0.003)
2<sup>nd</sup>: includes efficacy substudy (n = 365) and larger safety population (n = 2186) | 2<sup>nd</sup>: includes efficacy substudy (n = 365) and larger safety population (n = 2186)
† AEs with desvenlafaxine vs. tibolone and placebo (73.4% vs. 64.5% and 55.9%, respectively), most commonly nausea (31%), dizziness and constipation | 1<sup>st</sup>: AEs vs. placebo during week 1 only (p < 0.001), mostly nausea, dry mouth, constipation (but no diff. in BP)
‡ bleeding with tibolone vs. desvenlafaxine and placebo (23% vs. 12% (p = 0.024) and 9% (p = 0.001), respectively)
† discontinuations due to AEs during week 1 only vs. placebo (p < 0.001). Most commonly nausea (8.9%) and headache (3.8%)
Table 4 continued

| Clinical trial | Trial design and key participant BL data | Key outcomes | Results (p values vs. placebo) |
|----------------|-----------------------------------------|--------------|--------------------------------|
| 7a             | DBRCT: 3 US centres                     | 1*:          |                                |
| Joffe et al    | Mean age = 54.6 years                   | Mean daily # of HFs at weeks 4 and week 8 |                                |
| JAMA Internal Medicine (2014) [27] | Mean BMI = 28.3 kg/m² | HF severity at week 8 | 2*: baseline or % data not reported |
| Cann et al     | 339 randomised (2:2:3) to venlafaxine XR 75 mg/day (37.5 mg/day titrated to 75 mg/day over 1 week), oral 17-beta-oestradiol (ET) 0.5 mg/day or placebo for 8 weeks | MENQoL total and domain scores |                                                   |
| Menopause (2015) [28] | Venlafaxine followed by 2-week dose-tapering | Important AEs/SAEs |                                                   |
|                | 330 included in mITT analysis | Discontinuation due to AEs |                                                   |

**DBRCT** double-blind, randomised, placebo-controlled trial, **XR** extended release, **mod** moderate, **#** number, **HF** hot flushes/flashes, **FMS** vasomotor symptoms, ≧ greater than or equal to, ≦ less than or equal to, > greater than, < less than, **mg/d** milligrams/day, **(m)ITT** (modified) intention-to-treat, **BL** baseline, **1** primary, **2** secondary, **avg** average, **Δ change**, **sig** significant, **ns** not significant, ↑ increase, ↓ decrease, **diff** difference, **Tx** treatment, **(S)AEs** (serious) adverse events, **(S)** or (D) **BP** (systolic) or (diastolic) blood pressure, **LFTs** liver function tests, **ALT** alanine aminotransferase, **AST** aspartate aminotransferase, **NASH** non-alcoholic steatohepatitis, **ULN** upper limit of normal, **MENQoL** Menopause-Specific Quality of Life questionnaire, **HFRDIS** Hot Flash-Related Daily Interference Scale, **PEG** The Pain Enjoyment of Life and General Activity scale, **PHQ-9** 9-item Patient Health Questionnaire, **GAD-7** 7-item Generalized Anxiety Disorder questionnaire, **PSS** Perceived stress scale

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observed regarding generalisability and trial design which are important to note in order to place overall results in context and to consider when evaluating differences between treatment groups.

**Age**

HF symptoms are typically most severe around the final menstrual period. While mean age in the SNRI (53–55 years) and NK3Ra (54–55 years) trials was similar, the range was much greater in the SNRI trials (29–78 years vs. 41–65 years). This is relevant since it is unknown if treatment is equally effective in all menopause phases.

**Ethnicity**

Studies have shown ethnicity to affect HF prevalence with the highest frequency in Turkish women (97%) and the lowest in South American (47%) and Asian (45%) women [1]. Most SNRI trials predominately (~75–80%) recruited Caucasian women from North American centres which is a limitation since HF frequency can vary by population, often less in Asians, but greater in African-Americans [35]. NK3Ra trial participants were mainly Caucasian with few Asians, although an ongoing trial with fezolinetant is recruiting Asian women (NCT04234204) and a trial is planned in Chinese women (NCT04793204).

**Menopause Definition**

It is preferable to exclude perimenopausal women since symptoms might improve secondary to fluctuating hormone levels but be mistakenly attributed to the study drug. The
### Table 5: Neurokinin 3 receptor antagonists (NK3Ra) study results

| Clinical trial | Trial design and key participant BL data | Key outcomes | Results (p-values vs. placebo) |
|----------------|------------------------------------------|--------------|--------------------------------|
| 1a. Prague et al | DBRCT; Cross-over trial; 1 UK centre | | |
| The Lancet (2017) [29] | Mean age = 55 years (49–62) | Mean BMI = 25.85 kg/m² | |
| | 38 randomised to 4 weeks MLE4901 (40 mg BID) and 4 weeks placebo (BID) in random order separated by a 2-week washout period | 37 included in ITT analysis | |
| | | | 1a; # of weekly HFs during week |
| | | | 2a; HF severity, bother and interference scores at week 4 |
| | | | Important AEs/SAEs Discontinuations |
| | | | ↓ in weekly # of HFs from BL vs. placebo [ITT adjusted means: 19.35 (73%); 49.01 (28%); respectively; p < 0.001] |
| | | | ↓ HF severity score from BL vs. placebo [3.27 (44%); 5.70 (5%); p < 0.0001]; ↓ bother score (2.92 vs. 5.56, p < 0.0001), and ↓ interference score (7.94 vs. 26.48, p < 0.0001) |
| | | | Discontinuations (24%) higher than expected (mostly not MLE4901-related) |
| Ib | Prague et al. [30] reports a post hoc time course analysis of Prague et al. [29] to define therapeutic profile of MLE4901 by comparing the mean daily total of HFs at day 3, and mean weekly total after weeks 1, 2, 3, and 4 of both Tx periods, and also compared with week 2 of the BL period | Post hoc analysis of questionnaire data (minimum n = 33, maximum n = 35) | |
| | | | # of HFs |
| | | | HF severity, bother and interference |
| | | | Impact on sleep via: |
| | | | # of night-time HFs |
| | | | Individual MENQoL items |
| | | | Individual HFRDIS items |
| | | | ↓ # of HFs from BL vs. placebo by day 3 (72% vs. 21%); p < 0.0001), maintained through to week 4 |
| | | | ↓ HF severity, bother & interference continued to improve vs. placebo. At day 3: |
| | | | ↓ HF severity from BL by 38% (vs. 7%); p < 0.0001), which ↓ to −44% by week 4 (vs. 5%)
| | | | ↓ HF bother from BL by 39% (vs. 5%); p < 0.0001), which ↓ to −50% by week 4 |
| | | | ↓ HF interference from BL by 61% (vs. 24%); p = 0.0006), which ↓ to −70% by week 4 |
| | | | Impact of sleep: |
| | | | ↓ night-time HFs from BL vs. placebo at week 4 (78% vs. 22%; p < 0.0001). Improvements rapid, significant by day 3 vs. placebo (p < 0.0001)
| | | | Improved MENQoL psychosocial and physical domains at week 4. Authors suggested due to improved sleep since ‘difficulty sleeping’, ‘lethargy’ and ‘tiredness’ improved at week 4 (p < 0.0001, p = 0.00128, and p = 0.0002, respectively), and ‘lethargy’ and ‘tiredness’ improved by day 3 (p = 0.0474 and p = 0.0132, respectively), whereas ‘muscle ache’ and ‘physical strength’ did not improve |
| | | | Improved ‘sleep’ and ‘concentration’, significant by day 3 vs. placebo [sleep: p = 0.001; concentration: p = 0.0075] (HFRDIS) |

### 2

| Depypere et al | DBRCT: 8 Belgian centres | | |
| Journal of Clinical Endocrinology and Metabolism (2019) [31] | Mean age = 53.5 years (44–64) | Mean BMI = 25.8 kg/m² | |
| | 87 randomised (1:1) to fezolinetant 90 mg BID or placebo for 12 weeks | 87 included in mITT analysis (inferred from Fig. 2A but this is unclear) | |
| | | | Δ from BL in mean daily total VMS score (composite of # and severity) at week 12 |
| | | | Important AEs/SAEs Discontinuation due to AEs |
| | | | ↓ mean daily total VMS score from BL vs. placebo at week 4, 8 and 12 [week 12: 2.7 (91%); 14.4 (44%); all comparisons p < 0.0001] |
| | | | ↑ AEs considered Tx-related vs. placebo (30.2% vs. 25%), most commonly GI disorders (23% vs. 9%) |
| | | | ↑ discontinuations due to AEs (2.4%) vs. 0) |
| Clinical trial | Trial design and key participant BL data | Key outcomes | Results (p-values vs. placebo) |
|----------------|------------------------------------------|--------------|-------------------------------|
| 3a             | DBRCT; Dose-ranging 51 US centres       | 1*:           | All doses ↓ # of mod/severe HF from BL vs. placebo at week 4 (ranging 62–81%) vs. 59%) and week 12 (ranging 74–87%) vs. 55%); all p ≤ 0.024 |
| Fraser et al   | Mean age = 54.6 years (41–65)           | Mean Δ in # of mod/severe HF from BL at weeks 4 and 12 |                                  |
| VESTA          | Mean BMI = 28.4 kg/m²                    | Mean Δ in mod/severe HF severity from BL at weeks 4 and 12 |                                  |
| Menopause (2020) [32] | 356 randomised (1:1) to fezolinetant BID 15, 30, 60, or 90 mg or fezolinetant QD 30, 60, or 120 mg or placebo for 12 weeks | 2*: Important AEs/SAEs Discontinuation due to AEs | All doses ↓ mod/severe HF severity from BL vs. placebo at week 4 (ranging 29–54%) vs. 12%; all p ≤ 0.0322) and 60 mg BID, 90 mg BID, and 60 mg QD at week 12 ranging (52–53%) vs. 32%; all p ≤ 0.016 |
| 349 included in mITT analysis | | | |
| Santoro et al [33] reports results of 2′ endpoints from VESTA [32] | % achieving 50%, 70% and 90% ↓ in BL in # of mod/severe HF from BL | Responder analyses | 2′: AEs similar across Tx groups, with no indication of a dose effect. Most commonly nausea, diarrhoea, fatigue |
| Menopause (2020) [33] | | | |
| 4              | DBCRT; multiple-ascending-dose study; 3 US centres | Pre-specified exploratory efficacy endpoints: | 50 mg and 100 mg no significant ↓ in HF endpoints. (With 50 mg vs. placebo, mean HF frequency actually higher (p = 0.048) although median HF frequency no difference) |
| Trower et al   | Mean age = 55 years                     | Δ from BL in daily # of mod/severe HF at week 2 | ↓ # of mod/severe HF severity from BL at week 2 |
| RELENT-1       | Mean BMI = 28.18 kg/m²                  | Δ from BL in avg. daily HF severity at week 2 | Doses ≥ 150 mg improved symptoms early (within week 1 of Tx) |
| Menopause (2020) [34] | 76 randomised (3:1) to NT-814 vs. placebo within each of 4 sequential dose cohorts; 50, 100, 150, and 300 mg/day for 2 weeks | Δ from BL in daily # of waking due to night sweats | [HF severity vs. placebo with 150 mg [41%] vs. 13% (p < 0.001); 300 mg no sig. difference |
| 76 included in mITT analysis | Important AEs/SAEs Discontinuation due to AEs | | [waking due to night sweats vs. placebo [150 mg 81%] (p < 0.001), 300 mg 63% [p = 0.031] vs. 32%])]; AEs similar with placebo, 50 mg, 100 mg and 150 mg groups (slightly higher in 300 mg group) |
| Most common: mild somnolence and headache | | | Most discontinuations due to AEs vs. placebo: Numerical ↑ |
| No discontinuation due to AEs | | | |

DBRCT double-blind, randomised, placebo-controlled trial, XR extended release, mod moderate, # number, HF hot flushes/flashes, FMS vasomotor symptoms, ≥ greater than or equal to, ≤ less than or equal to, > greater than, < less than, mg/d milligrams/day, (m)ITT (modified) intention-to-treat, BL baseline, 1° primary, 2° secondary, avg average, Δ change, sig significant, ns not significant, ↑ increase, ↓ reduced, diff difference, Tx treatment, (S)AE (serious) adverse events, (S) or (D) BP (systolic) or (diastolic) blood pressure, LFT liver function tests, ALT alanine aminotransferase, AST aspartate aminotransferase, NASH non-alcoholic steatohepatitis, ULN upper limit of normal, MENQoL Menopause-Specific Quality of Life questionnaire, HFRDIS Hot Flashes-Related Daily Interference Scale, PEG The Pain Enjoyment of Life and General Activity scale, PHQ-9 9-item Patient Health Questionnaire, GAD-7 7-item Generalized Anxiety Disorder questionnaire, PSS Perceived stress scale.
FDA [36] advise using the following menopause definition: 12 months spontaneous amenorrhea, or 6 months amenorrhea plus FSH > 40mIU/ml, or 6 weeks post-surgical menopause. Two SNRI trials did not appear to fulfil the FDA definition and thus could have included perimenopausal women [20, 24]. All NK3Ra trials fulfilled the FDA definition, apart from Depypere et al. [31] who allowed amenorrhea ≥ 3 months if FSH > 40 IU/L and oestradiol < 0.21 nmol/L (which usually indicates premature ovarian insufficiency but oestrogen levels can vary in early stages).

**HF Frequency and Severity**
The FDA [36] advise enrolling subjects with ≥ 7 moderate-severe HFs/day (or ≥ 50/week). All trials fulfilled this except 2 SNRI trials [20, 27] which only required ≥ 14 HFs/week and not all were moderate or severe. Joffe et al. [27] reasoned that by not requiring ≥ 7 HFs/day (which only occurs in 7–9%), they enrolled a more generalisable population, many of whom do seek HF treatment. However, mild HFs may be less challenging to relieve and thus Joffe’s efficacy data could be potentially exaggerated.

**HF Stability**
A stable HF pattern is needed to minimise the risk of a spontaneous HF reduction being attributed to a study drug. Most trials required a consistent minimum or < 50% change in pattern over 1–2 baseline weeks. However, two SNRI trials [20, 21] failed to assess baseline stability and hence their results are potentially unreliable. One SNRI trial [23] and one NK3Ra trial [32] required ≥ 50 HFs over any 7 consecutive days during the 35-day screening period, yet separately used the week prior to randomisation as the baseline comparator, potentially underestimating baseline HF comparisons.

**Baseline Characteristics**
Treatment and control groups were generally well balanced. However, Evans et al. [20] reported higher alcohol consumption (a potential HF trigger) in their SNRI group but addressed this using sensitivity analysis. Among NK3Ra trials, Trower et al. [34] reported that the NT-814 50 mg group had a higher baseline HF frequency, severity and night-time awakenings which likely accounts for the unexpected observation of 50 mg showing less reduction in HFs compared to placebo.

**Comorbidity**
All trials recruited ‘healthy’ women. While safety data for the use of SNRIs in complex patients has been collected elsewhere, it will be important to establish efficacy and safety of NK3Ras in complex patients during phase 3 or post-market registry settings.

**Washout of Prior HF Treatment**
To avoid potential confounding effects, the FDA [36] advises different HRT washout periods depending on administration route. However, a washout was not specified in one SNRI trial [20], and too short a washout was used for one SNRI [27] and one NK3Ra trial [31], thus a spill-over effect may have occurred.

**Placebo Control**
The inclusion of a placebo arm comparator was validated after a marked reduction in HF frequency was noted after placebo administration in trials [SNRI range (across all 7 trials) 25–58%; NK3Ra range 28–55%]. One method to attenuate a large placebo response may be to specify increased HF bother or a similar lifestyle measure as an inclusion criterion. Pinkerton et al. [25], for example, which had the lowest placebo response at week 12, required a score of ≥ 12 on the Greene Climacteric Scale (a standard questionnaire assessing physical and psychological menopausal symptoms) at study entry.

**Power and Sample Size**
All SNRI trials undertook a power calculation, except Evans et al. [20] which may have been underpowered as the reduction in HF frequency of 1.4/day was not statistically significant (p = 0.06), despite a statistically significant reduction in patient-perceived severity score. Initial NK3Ra trials lacked prior efficacy data to estimate treatment effect but instead used power calculations based on the ability to detect a treatment effect double the...
anticipated ~ 25% placebo effect [29] or based on previous HF diary trials [31]. Fraser et al. [32] powered based on prior NK3Ra efficacy [31], but, due to drop-out, failed to achieve the planned 40 subjects in 7/8 groups which may have affected 12-week HF severity evaluation (in which only 3 groups showed a significant reduction). Trower et al. [34] based sample size on a previous pilot study suggesting 8 subjects per group was adequate, but since the 150 mg group achieved all primary endpoints, the failure of 300 mg to show a statistically significant reduction in HF severity at week 12 may have been due to being underpowered.

**HF Recording**
All but three trials used retrospective paper diaries to assess HF frequency and severity. However, errors in compliance are major sources of bias and backfilling is common [37]. Prospective time-stamped electronic diaries may give superior timeliness and completion versus paper diaries [37] and were used in three trials [29, 31, 32]. Encouraging respondents to report HFs as they occur (as in one NK3Ra trial [29]), rather than at the end of the day/the following morning, also reduces recall bias.

**Estimating Night Sweats**
Use of the ‘number of night-time awakenings’ in SNRI [21–23] and NK3RAs trials [34] is problematic. Night-sweats may not produce full awakenings and may be under-reported. Disparity between subjective and objective reporting widens at night [38]. Thus, questionnaires such as MENQoL, which evaluate sleep quality [28, 30, 31, 33] may be useful since improved scores in ‘concentration’, ‘difficulty sleeping’, ‘tiredness’ and ‘lethargy’ correlate with HF therapeutic benefit and may more accurately reflect the impact on participants’ lives.

**Discontinuation**
If patient drop-out is high, statistical power may be lost. With SNRI trials, early discontinuation due to AEs, often due to nausea, was common (5–29%), especially during week 1. When treating depression, SNRIs are usually titrated over 1–2 months to reduce AEs [39]. It is surprising that only four of seven SNRI trials [20, 22, 25, 27] used titration and, even in those trials that did, the titration was very rapid (over 1 week) [22, 27]. In NK3Ra trials, early discontinuation due to AEs was uncommon (5–7%). Since HFs can continue for 12 years [4], longer trials with SNRIs and NK3RAs are warranted to evaluate long-term efficacy and tolerability/safety.

**SNRI Tolerability and Safety**
Most SNRI trials were short (< 12 weeks), apart from two trials which extended to 26 and 52 weeks, albeit with high dropout rates [23, 26]. Higher-dose SNRIs have been associated with treatment-emergent hypertension (likely due to increased potentiation of noradrenergic neurotransmission). A significant or numerical increase in blood pressure was reported in three SNRI trials [21, 22, 27], although none was powered to evaluate this endpoint (Table 4). A 1-year CV safety study among 2118 subjects with HFs [40] and pooled data from >6000 subjects receiving desvenlafaxine for various indications [21] did not report a significant excess of CV events but 1 year may be too short; the Framingham Heart Study showed an association between ~ 10 mmHg increase in blood pressure and increased CV events took 4–6 years to emerge [41].

**NK3Ra Tolerability and Safety**
Since only short-duration phase 2 data have been published, it was not possible to fully evaluate NK3Ra tolerability and safety. In two trials, participants taking NK3RAs reported a higher frequency of minor GI disorders [31, 32], which could be related to NK3R expression in the GI tract [42]. However, the AE incidence did not appear dose-dependent. Somnolence was also more common (69% vs. 17%) with the highest (300 mg) dose of NT-814 [34] likely due to its additional NK1Ra action [43]. Interestingly, this could be an advantage for women with sleep disturbance. Asymptomatic rises in transaminases >3 times the upper limit of normal were reported with MLE4901 [29] and fezolinetant [32] but bilirubin did not rise >2
times upper limit of normal. Liver function tests (LFTs) returned to normal after drug discontinuation. However, further safety analyses are required.

**Potential Study Bias**

Overall risk of bias was low (Table 3). Selection bias was minimised by randomised study designs. Unreported data leading to potential reporting bias was noted, including pre-specified data on quality of life [21–23, 33], 50% HF responder rates [21] and apparent differences in gastrointestinal AE frequency in abstract versus table [31]. Potential conflicts of interest were considered under reporting bias but most trials declared conflicts and often had industry co-authors, which is common for these types of studies. All trials were double-blind (except one [20] using matched placebo) which should have reduced performance bias and detection bias, although this is difficult to fully exclude. One SNRI trial included an HRT arm [24] in which subjects experienced significant bleeding compared with the desvenlafaxine or placebo groups which likely unblinded treatment allocation. High drop rates of ≥20% occurred in several trials [20, 21, 23, 24, 26, 29] leading to potential attrition bias [44], although this risk was reduced by appropriate use of intention-to-treat (ITT) or modified ITT rather than per-protocol analysis for primary outcomes.

**Potential Additional Benefits of NK3Ra**

**Onset of Therapeutic Effect**

Prague et al. [30] reported symptom improvement after ~48 h with NK3Ras, with maximum effect by day 3. Thus, NK3Ras would be expected to give earlier symptom relief versus SNRIs, especially given the slow dose titration recommended for SNRIs.

**Sleep Quality/Concentration**

Improved sleep ± concentration was noted in three NK3Ra trials [30, 31, 34]. This is likely attributed to multiple factors including reduced sleep disruption from HFs, and the attenuation of NK3R actions in melanin-concentrating hormone neurons (involved in the sleep–wake cycle) and in the prefrontal cortex, an important area for concentration [45], whereas the additional NK1Ra action of NT-814 may attenuate substance P-induced arousal and facilitate sleep [43]. In contrast, SNRIs are non-sedating [46], hence a reduction in night-time awakening with SNRIs is likely due to HF reduction and anxiolysis.

**Mood**

SNRIs improve depressed mood (another common menopausal symptom). Although it is unclear if NK3Ras directly affect mood, the NK1Ra action of NT-814 may be of benefit [47]. In addition to pharmacotherapy, it may also be worth considering cognitive behavioural therapy (CBT) if mood is a particularly limiting symptom, as the positive effects of CBT appear to be sustained over time [48].

**CV Safety**

After menopause, CV risk increases. In contrast to SNRIs which may be associated with hypertension leading to increased CV risk (discussed above), NK3Ras in rats reversed spontaneous hypertension and lowered heart rate [49] via reducing midbrain dopaminergic signalling in the ventral tegmental area that highly expresses NK3Rs [50]. Vasopressin neurones also express NK3Rs [51], and NKB activity is potentiated by thromboxane A2 [52] which might also represent useful therapeutic targets for NK3Ras.

**Limitations of This Analysis**

Conclusions of our qualitative analysis are limited by published data for NK3Ra being only phase 2 trials. Longer duration phase 3 trials are ongoing, and findings will provide further insights into efficacy and safety outcomes [53]. Availability of phase 3 NK3Ra data would also facilitate meta-analysis, enabling determination of a weighted pooled estimate for HF reduction in NK3Ra versus SNRI trials, although the ideal trial to compare NK3Ras versus SNRIs for reduction of menopausal HFs would be a large phase 3 head-to-head trial.
CONCLUSIONS

Menopausal HFs can significantly impact a woman’s quality of life, but some women cannot or do not wish to take HRT to alleviate their symptoms. Current non-hormonal options are suboptimal due to variable efficacy and low tolerability. The recent discovery that NKB-NK3R signalling is implicated in the generation of menopausal HFs has led to recent clinical trials using NK3Ras. Qualitative analysis of these trials indicates that NK3Ras lead to greater reductions in mean HF frequency, HF severity and night-sweats with good short-term tolerability compared to SNRIs. Efficacy and safety data (including careful evaluation of LFTs) from phase 3 trials are awaited with interest.

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