Deleterious Effects of Cold Air Inhalation on Coronary Physiological Indices in Patients With Obstructive Coronary Artery Disease

Rupert P. Williams, PhD; Kaleb N. Asrress, PhD; Matthew Lumley, PhD; Satpal Arri, BSc; Tiffany Patterson, PhD; Howard Ellis, BSc; Vasiliki Manou-Stathopoulou, BSc; Catherine Macfarlane, BSc; Shruthi Chandran, BSc; Kostantinos Moschonas, MRes; Pippa Oakeshott, MD; Timothy Lockie, PhD; Amedeo Chiribiri, PhD; Brian Clapp, PhD; Divaka Perera, PhD; Sven Plein, PhD; Michael S. Marber, PhD; Simon R. Redwood, MD

Background—Cold air inhalation during exercise increases cardiac mortality, but the pathophysiology is unclear. During cold and exercise, dual-sensor intracoronary wires measured coronary microvascular resistance (MVR) and blood flow velocity (CFB), and cardiac magnetic resonance measured subendocardial perfusion.

Methods and Results—Forty-two patients (62±9 years) undergoing cardiac catheterization, 32 with obstructive coronary stenoses and 10 without, performed either (1) 5 minutes of cold air inhalation (5°F) or (2) two 5-minute supine-cycling periods: 1 at room temperature and 1 during cold air inhalation (5°F) (randomized order). We compared rest and peak stress MVR, CFB, and subendocardial perfusion measurements. In patients with unobstructed coronary arteries (n=10), cold air inhalation at rest decreased MVR by 6% (P=0.41), increasing CFB by 20% (P<0.01). However, in patients with obstructive stenoses (n=10), cold air inhalation at rest increased MVR by 17% (P<0.01), reducing CFB by 3% (P=0.85). Consequently, in patients with obstructive stenoses undergoing the cardiac magnetic resonance protocol (n=10), cold air inhalation reduced subendocardial perfusion (P<0.05). Only patients with obstructive stenoses performed this protocol (n=12). Cycling at room temperature decreased MVR by 29% (P<0.001) and increased CFB by 61% (P<0.001). However, cold air inhalation during cycling blunted these adaptations in MVR (P=0.12) and CFB (P<0.05), an effect attributable to defective early diastolic CBF acceleration (P<0.05) and associated with greater ST-segment depression (P<0.05).

Conclusions—in patients with obstructive coronary stenoses, cold air inhalation causes deleterious changes in MVR and CFB. These diminish or abolish the normal adaptations during exertion that ordinarily match myocardial blood supply to demand. (J Am Heart Assoc. 2018;7:e008837. DOI: 10.1161/JAHA.118.008837.)

Key Words: cold • coronary • coronary flow • coronary microvascular resistance • physiology • wave intensity analysis

Cold air inhalation during vigorous physical exertion, especially shoveling snow,1 results in significant increases in exertion-related cardiac deaths.2 Cold air alone is associated with an increased incidence of myocardial infarction and cardiac mortality,3 independent of confounding factors such as respiratory illness.3 Exertion in the cold has also been shown to reduce time to angina, and time to electrocardiographic ST-segment depression,4 compared with exertion at room temperature.

The pathophysiology of sudden cardiac death during vigorous exertion in cold air is unclear but likely to be multifactorial.4–7 One of the triggering events may be the deleterious effect of cold air on coronary blood flow.5,8 Patients with obstructive coronary stenoses have less coronary vasodilator reserve and maybe particularly vulnerable to such deleterious effects.9 Changes in coronary blood flow are principally mediated by changes in coronary microvascular resistance (MVR). However, to date no in vivo MVR measurements have been taken during cold air inhalation.

Dual-sensor intracoronary guidewires now allow accurate continuous in vivo measurement of distal coronary artery...
Clinical Perspective

What Is New?

- This study was performed to further our understanding of the pathophysiological effects of exertion in the cold.
- We used cold air inhalation as a physiological cold stressor and designed a study protocol enabling supine exercise to be performed on the cardiac catheter laboratory table during coronary angiography.
- Additionally, we used dual-sensor pressure and flow intracoronary guidewires to enable continuous and simultaneous acquisition of coronary blood flow and microvascular resistance.
- We demonstrated that cold air inhalation during supine exercise severely blunted coronary vasodilation and the increase in coronary blood flow compared with supine exercise at room temperature, significantly increasing myocardial ischemia.

What Are the Clinical Implications?

- Our study demonstrates the potential hazardous mismatch between coronary flow and myocardial work caused by even brief periods of cold air inhalation with and without exercise, particularly in patients who have coronary artery disease and who have a reduced vasodilator reserve.
- Doctors should inform patients with obstructive coronary stenoses (1) to avoid exertion in cold air where possible, or at least to wrap up warmly (ideally involving coverage of the cheeks, mouth, and forehead); (2) that their exercise tolerance may be less in cold air; and (3) indoor warm-up exercise before exertion in cold air might attenuate these deleterious effects.

Methods

The data, analytic methods, and study materials have been made available to other researchers for purposes of reproducing the results or replicating the procedure.7

Between April 2013 and July 2015, we recruited 42 patients (62±9 years) from routine coronary angiography waiting lists, in 2 distinct groups (Figure 1):

1. Patients with obstructive coronary stenoses (n=32): defined by either a coronary artery stenosis >50% on quantitative coronary angiography15 or fractional flow reserve ≤0.8.16 These patients had typical anginal symptoms and multiple risk factors, or known obstructive coronary artery stenoses awaiting percutaneous coronary intervention.
2. Patients with unobstructed coronary arteries (n=10): defined by a coronary artery stenosis <30% on quantitative coronary angiography. The majority of these patients had atypical chest pain with some cardiac risk factors, and some had a mildly positive or equivocal exercise treadmill test or ischemia test.

Exclusion criteria for all groups were as follows: patients with 30% to 50% stenosis on quantitative coronary angiography; unstable symptoms; impaired left ventricular function; severe chronic obstructive pulmonary disease or asthma (to minimize risk of bronchospasm with cold air inhalation); and an inability to undertake exercise and standard contraindications to CMR.

All vasoactive medications were stopped 48 hours before the procedure. Subjects gave written informed consent in accordance with the protocol approved by the institutional ethics review committee (REC 13/LO/0759). The study was registered with the National Institute for Health Research (NIHR) UK Clinical Research Network (UKCRN) portfolio database (ID 131491).

Study Interventions

Cold air inhalation

Cold air was delivered via a loose-fitting facemask attached to the patient’s face for 5 minutes, at a fixed temperature of 5°F and a flow rate of 50 L/min (using an adapted cryotherapy device with a closed cooling circuit: Cryo6 Cold Air Device; Zimmer Medizin Systeme, Ulm, Germany). The cold air chilled the cheeks, nose, mouth, and laryngeal cold receptors.5

Exercise

An adapted supine ergometer (Ergosana, Schiller, Germany) was attached to the cardiac catheterization laboratory table.17

pressure and Doppler-flow velocity, permitting calculation of MVR.10 Continuous acquisition of these data also enables application of wave intensity analysis to identify the primary events driving coronary blood flow and myocardial work caused by even brief periods of cold air inhalation with and without exercise, particularly in patients who have coronary artery disease and who have a reduced vasodilator reserve.

The aims of this descriptive study are as follows:

1. To determine the effect of the presence or absence of an obstructive coronary artery stenosis on the hemodynamic response (MVR, CBF, and wave intensity analysis) to a period of cold air inhalation at rest.
2. In patients with an obstructive coronary artery stenosis, to determine the additional effect of cold air inhalation during a period of exercise on the same hemodynamic variables, compared with exercise during room temperature as a control.
Exercise started at 50 W, with 20-W increments every minute. It continued for 5 minutes unless the following occurred: severe angina, breathlessness, fatigue, or target heart rate achieved (220 beats per minute—aige).

### Study Design

All studies were performed at room temperature (70°F). The potential psychological effect of wearing the facemask at
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baseline (without cold air inhalation) was assessed in all patients. Patients were randomly assigned to complete 1 of the 2 protocols (Figure 1), whereby baseline and peak stress measurements were acquired before and after either:

Protocol 1. Cold air inhalation at rest (n=30):
5 minutes of cold air inhalation at rest, in either an (1) invasive or (2) noninvasive protocol, as below:

a. Cardiac Catheterization Laboratory Protocol (invasive coronary flow velocity and pressure data): in patients with obstructed coronary stenoses (n=10) and patients with unobstructed coronary arteries (n=10).
b. CMR Protocol (subendocardial to subepicardial perfusion gradients): in a further separate group of patients with obstructive coronary stenoses (n=10).

OR

Protocol 2. Exercise with and without cold air inhalation (n=12):
Two 5-minute supine-cycling exercise periods: 1 at room temperature AND 1 during cold air inhalation. Only patients with obstructive coronary stenoses (n=12) were recruited to this study, and only the Cardiac Catheterization Laboratory Protocol was used. Exercise in room temperature was used as our control. The exercise periods were performed in randomized order to minimize potential warm-up effects of prior exercise.17 There was a minimum of 15 minutes rest between exercise periods to allow hemodynamics to return to baseline.

Study Protocols

Cardiac catheterization laboratory protocol

Patients were catheterized via the right radial or femoral artery with standard 6-Fr coronary guide catheters (although radial arterial access was mandated in all patients undergoing exercise protocols) (Video S1, Figure S1). Patients received 300 mg of aspirin and 600 mg of clopidogrel orally, and 70 units/kg heparin intra-arterially. A dual pressure and Doppler sensor-tipped 0.014-in intracoronary wire (ComboWire, Volcano Corp, San Diego, CA) was used to measure coronary pressure and Doppler-flow velocity. Wire pressure was first equalized to that recorded by the fluid-filled manometer and then the wire tip was advanced into the distal target coronary artery and manipulated until a stable and optimal Doppler-flow velocity trace was obtained. Hemodynamic measurements were taken under resting conditions and continuously during each stressor.

CMR protocol

Cine and myocardial perfusion images were acquired on a 3-T CMR scanner (Achieva Tx, Philips Medical Systems, Best, the Netherlands) at baseline and during peak cold air inhalation at rest (scan started 3 minutes after cold air inhalation commenced). High spatial resolution (1.2 x 1.2 mm) perfusion images were acquired in midsystole using a saturation recovery gradient echo method.12,13 Further details are specified in Data S1.

Hemodynamic Analyses

Aortic pressure, distal coronary artery pressure, and average peak Doppler-flow velocity data were sampled at 200 Hz and stored on disk for off-line analysis. Data were imported into custom software (Study Manager, Academic Medical Center, University of Amsterdam, the Netherlands), and 10 consecutive beats were extracted for analysis. Another custom-made program, Cardiac Waves (Kings College London, UK), was then used to perform all further analyses. More detailed methods are specified in Figure S1.

Average distal coronary artery Doppler-flow velocity (CBF) and pressure (Pd) were obtained from the ComboWire. MVR was calculated as (Pd/CBF).10 “Minimal” MVR was also calculated as previously defined,18 through identification of the wave-free period in diastole with wave intensity analysis when resistance is at its lowest. Minimal MVR therefore excludes systolic compressive resistance and better informs vascular resistance, and has been shown not to be influenced by coronary artery stenosis severity.18 Trans-stenotic pressure gradient (aortic pressure (Pa)−Pd), Pd/Pa ratio, and epicardial stenosis resistance (Pd−Pa/average peak velocity)19 were also calculated as measures of stenosis severity. Epicardial stenosis resistance, defined as the ratio of the pressure drop across the stenosis to distal coronary flow velocity, normalizes the pressure drop for the magnitude of epicardial coronary flow velocity at which it was obtained, providing a more objective assessment of hemodynamic stenosis severity. Moreover, these data combined allow the selective evaluation of epicardial and MVR to coronary blood flow.20

Coronary wave intensity analysis: Continuous acquisition of simultaneous changes in distal CBF and pressure also allows insight into the phasic components of forces driving CBF. Net coronary wave intensity (di) was calculated as the product of the derivatives of ensemble-averaged distal coronary pressure (Pd) and Doppler-flow velocity (CBF), so that: di=dPd/dt×dCBF/dt.21 We calculated, as previously described,22,23 the 2 dominant waves that drive coronary perfusion (Figure 2):
(1) The forward compression wave, a systolic acceleratory wave generated by sudden increases in aortic pressure at the inlet during ventricular ejection. (2) The backward expansion wave, an early diastolic acceleratory wave generated by microcirculatory suction from the rapid reduction in left ventricular pressure during ventricular relaxation (relieving compression of intramyocardial vessels).22,23

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Aortic pulse wave analysis (from central arterial pressure waveforms): Augmentation index (a central measure of afterload), rate pressure product (a measure of myocardial oxygen consumption), and diastolic time fraction were calculated as previously described.17

Myocardial perfusion analysis: Myocardial perfusion CMR images were analyzed using dedicated software (EasyScil prototyping [Philips Medical Systems, Best, the Netherlands] and Matlabatory [MathWorks Inc, Natick, MA]). Absolute myocardial blood flow and subendocardial to subepicardial perfusion gradients were calculated as previously described.12,13 These maximize the high spatial and temporal resolution of the CMR perfusion sequence, and present the distribution of myocardial blood flow between the subendocardium and subepicardium over time12,13 (Figure S2). An increased gradient suggests subendocardial ischemia, using prospectively established thresholds.12 Further details are specified in Data S1.

Statistical Methods

Continuous variables were tested for normality using the Shapiro–Wilk test and presented as mean±SEM when data were normally distributed or as median with interquartile range.

All data were analyzed (1) at baseline and (2) during peak stress. Continuous data were examined every 30 s throughout the invasive protocol and peak stress was defined as the data set with the highest rate pressure product. A repeated-measures 2-way ANOVA was performed with matched values stacked into a subcolumn. To assess differences between rest and stress data, Sidak’s multiple comparison test was then performed (confidence intervals and significance were computed, and an adjusted significance threshold of $P \leq 0.05$ was used for these comparisons). Between-group differences in absolute change were assessed using paired or unpaired t tests (continuous and normal data) or Mann–Whitney U test or Wilcoxon signed-rank test (non-normal continuous data) as appropriate. A 2-tailed test for significance was performed in all of the analyses.

Investigators performing data analyses were blinded to all clinical data. Statistical analyses were performed using GraphPad Prism 6.0 (GraphPad, San Diego, CA). $P$ values of $\leq 0.05$ were considered statistically significant.

Results

Study Population

Of the 58 patients consented into the study across the different protocols, Figure 1 shows that 42 successfully completed a full protocol (32 completed a cardiac catheterization laboratory protocol and 10 completed a CMR protocol). Sixteen patients were excluded: 14 from the cardiac catheterization laboratory protocol and 2 from the CMR protocol. In addition, 6 data sets were excluded from coronary hemodynamic and wave intensity analyses because of poor quality Doppler-flow waveforms. Table 1 shows patient characteristics. Wearing the facemask

![Wave Intensity](image.png)

**Figure 2.** Coronary wave intensity (WIA) profile at rest. This figure is a typical example of a WIA profile obtained at rest from a patient with unobstructed coronary arteries. The 2 shaded areas represent the forward compression and backward expansion wave intensities.
(without cold air inhalation) at baseline did not cause significant changes to any variables.

**Coronary MVR and CBF**

**Cold air inhalation at rest**

At baseline before cold air was applied, patients with obstructive stenoses had significantly lower minimal MVR than patients with unobstructed coronary arteries \( P < 0.05 \) (Figures 3A and 4, Table 2). In patients with unobstructed coronary arteries, cold air inhalation at rest caused 6% and 8% reductions in MVR \( (P=0.41) \) and minimal MVR \( (P=0.27) \), respectively. This enabled a 20% increase in CBF \( (P<0.01) \). However, in patients with obstructive coronary stenoses, there were deleterious 17% and 24% increases in MVR \( (P<0.01) \) and minimal MVR \( (P<0.01) \), respectively, resulting in a 3% reduction in CBF \( (P=0.85) \). This resulted in between-group differences in absolute change for both MVR \( (P=0.005) \) and CBF \( (P=0.006) \).

**Exercise with and without cold air inhalation in patients with obstructive stenoses**

During exercise at room temperature, MVR and minimal MVR decreased by 29% \( (P<0.001) \) and 41% \( (P<0.001) \), respectively, associated with a 61% increase in CBF \( (P<0.001) \) (Figures 5A and 6, Table 2). However, MVR and minimal MVR decreased by only 13% \( (P=0.12) \) and 12% \( (P=0.48) \), respectively, during exercise in cold air inhalation, associated with a smaller (35%) increase in CBF \( (P<0.05) \). Paired group analysis also demonstrated that absolute reductions in MVR and minimal MVR during exercise with cold air inhalation were significantly smaller than during exercise in room temperature \( (P=0.04, \text{Figure 5A}) \) and \( (P<0.01) \), respectively.

**Coronary Wave Intensity Analysis**

**Cold air inhalation at rest**

In patients with unobstructed coronary arteries, cold air inhalation increased the energy of the 2 dominant waves that drive coronary perfusion: the forward compression wave by 98% \( (P<0.001) \) and the backward expansion wave by 71% \( (P<0.001) \) (Figure 3B, Table 2). However, in patients with obstructive coronary stenoses, cold air inhalation actually caused a 5% reduction in the magnitude of the forward compression wave \( (P=0.93) \) and abrogated the backward expansion wave response to a 12% increase \( (P=0.79) \). This resulted in between-group differences in absolute change for both the forward compression wave \( (P=0.001) \) and the backward expansion wave \( (P=0.02) \) (Figure 3B).

**Exercise with and without cold air inhalation in patients with obstructive stenoses**

Exercise at room temperature resulted in profound increases to the forward compression and backward expansion waves of 208% \( (P<0.05) \) and 239% \( (P<0.001) \), respectively (Figures 5B and 6, Table 2). However, during exercise in cold air inhalation, the backward expansion wave only increased by 95% \( (P=0.13) \). Paired group analysis also demonstrated that the absolute increases in the backward expansion wave during exercise with versus without cold air inhalation were

**Table 1. Patient Characteristics**

| Stressed (s)                     | Cold Air at Rest: Cath Lab Study | Cold Air at Rest: Cath Lab Study | Cold Air at Rest: CMR Study | Exercise With and Without Cold Air: Cath Lab Study |
|----------------------------------|----------------------------------|----------------------------------|----------------------------|---------------------------------------------------|
| Patient Demographic              | Unobstructed Coronaries          | Obstructive Stenoses             | Obstructive Stenoses       | Obstructive Stenoses                               |
| Number of patients               | 10                               | 10                               | 10                         | 12                                                |
| Age, y                           | 67.3±2.9                         | 64.6±2.6                         | 59.9±2.2                   | 58.8±2.7                                          |
| Male                             | 6 (60%)                          | 8 (80%)                          | 8 (80%)                    | 12 (100%)                                         |
| BMI, kg/m²                       | 29.1±1.3                         | 33.9±2.3                         | 29.9±1.7                   | 30.2±2.0                                          |
| Medical history                  |                                  |                                  |                            |                                                   |
| Hypertension                     | 9 (90%)                          | 9 (90%)                          | 7 (70%)                    | 9 (69%)                                           |
| Diabetes mellitus                | 2 (20%)                          | 4 (40%)                          | 4 (40%)                    | 2 (15%)                                           |
| Hypercholesterolemia             | 9 (90%)                          | 9 (90%)                          | 8 (80%)                    | 11 (85%)                                          |
| Smokers                          | 3 (30%)                          | 6 (60%)                          | 7 (70%)                    | 7 (54%)                                           |
| Procedural details               |                                  |                                  |                            |                                                   |
| Stenosis of target lesion, %     | 19.9±2.4                         | 76.5±4.1                         | 72.9±3.8                   | 68.6±4.7                                          |
| Target vessel (LAD/Cx/RCA)       | 7/2/1                            | 7/1/2                            | 3/4/3                      | 10/2/1                                            |

Values are mean±SEM or %. BMI indicates body mass index; Cath lab, cardiac catheterization laboratory; CMR, cardiac magnetic resonance; Cx, circumflex artery; LAD, left anterior descending artery; Obstructive stenoses, patients with obstructive coronary stenoses; RCA, right coronary artery; unobstructed coronaries, patients with unobstructed coronary arteries.
**Figure 3.** Cold air inhalation at rest. Change from baseline in patients with and without obstructive coronary stenoses, with the results subcategorized as per the following: (A) coronary hemodynamic results, (B) coronary wave intensity analysis results, and (C) aortic pulse wave analysis results.
without obstructive coronary stenoses (Figure 3C, Table 2). However, diastolic time fraction was deleteriously decreased in patients with obstructive coronary stenoses ($P<0.04$), but not in patients with unobstructed coronary arteries ($P=0.49$).

**Exercise with and without cold air inhalation in patients with obstructive stenoses**

Increases in rate pressure product were greater during exercise with versus without cold air inhalation, although this did not reach significance ($P=0.06$) (Figures 5C and 6, Table 2). Exercise at room temperature decreased augmentation index by 47% ($P<0.05$), but the addition of cold air inhalation during exercise abolished this adaptive reduction to only 5% ($P=0.95$). The diastolic time fraction reduced during exercise with and without cold air inhalation (by 31% and 21%, respectively, both $P<0.001$). However, overall, the addition of cold air inhalation during exercise caused a further deleterious absolute reduction in diastolic time fraction compared with exercise in room temperature ($P=0.05$).

**Electrocardiographic ST-Segment Depression**

Apart from 1 patient with an obstructive stenosis who withdrew after developing angina, cold air inhalation at rest did not result in any reported angina or measurable ST-segment depression. Maximal ST-segment depression was greater during exercise with cold air inhalation versus without (0.17±0.02 mV versus 0.12±0.02 mV, $P=0.02$). The maximum tolerated exercise duration was 313±15 s versus 337±24 s for exercise with versus without cold air ($P=0.13$), with similar workloads (116±7 W versus 115±7 W, respectively, $P=0.84$). This is equivalent to between 6.5 and 9 metabolic equivalents or stage 2-3 Bruce protocol.

**Stenosis Severity**

Measures of stenosis severity did not significantly increase in either group during cold air inhalation at rest (Table 2), although there was a trend ($P=0.11$) towards reduced epicardial stenosis resistance in patients with unobstructed coronary arteries and increased epicardial stenosis resistance in patients with obstructive stenoses. During exercise at room temperature, there were significant increases in trans-stenotic gradient ($P<0.001$), $P_{a}/P_{s}$ ratio ($P<0.01$), and epicardial stenosis resistance ($P<0.01$). Unexpectedly during exercise in cold air, there were smaller increases seen in these parameters, and significantly lower peak trans-stenotic gradient and $P_{a}/P_{s}$ ratio ($P=0.02$ and $<0.01$, respectively).

**CMR Analysis**

Ten separate patients with obstructive coronary stenoses underwent the CMR protocol (Figure 2, Table 1). Cold air
inhalation at rest caused significant deleterious increases in subendocardial to subepicardial transmural perfusion gradients from baseline, suggesting inadequate subendocardial perfusion (Figure S3). In keeping with invasive CBF data, there was no associated substantial increase in overall myocardial blood flow during cold air inhalation (0.92±0.14 mL/g per minute versus 1.07±0.11 mL/g per minute, P=0.19).

**Discussion**

For the first time, we have combined intracoronary measurements of pressure and Doppler-flow during cardiac catheterization to understand the pathophysiology of cold air inhalation during exercise. These measurements show that in patients with obstructive coronary stenoses, cold disrupts the fundamental physiological adaptations that ensure myocardial blood supply matches demand during exercise. In these patients, cold air inhalation prevented the decrease in myocardial MVR needed to increase blood flow during exercise, while at the same time increasing cardiac work through heightened afterload (Figure 6). High-resolution myocardial perfusion CMR demonstrated that the increases in myocardial MVR and cardiac afterload manifested as a cold-induced deficit in subendocardial perfusion. These deleterious changes render the myocardium vulnerable to ischemia, infarction, and potentially fatal cardiac arrhythmias.

**Table 2. Catheter Laboratory Coronary and Aortic Hemodynamics and Wave Intensity Analysis**

| Condition                          | Cold Air Inhalation at Rest | Exercise at Room Temp | Exercise During Cold Air |
|------------------------------------|-----------------------------|-----------------------|--------------------------|
| **Patient Demographic**            | Unobstructed Coronaries     | Obstructive Stenoses  | Obstructive Stenoses     | Obstructive Stenoses     |
| Patient Demographic                | Baseline                    | Peak                  | Baseline                 | Peak                     |
| **Stenosis severity**              |                             |                       |                          |                          |
| Pd (mean)/Pa (mean) ratio          | 0.96±0.01                   | 0.97±0.01             | 0.83±0.08                | 0.88±0.06                | 0.82±0.05*               | 0.90±0.04               | 0.87±0.05*†             |
| Trans-stenotic gradient, mm Hg     | 4.2±0.9                     | 3.4±1.3               | 15.9±7.2                | 19.0±9.3                | 14.2±5.9                | 24.8±6.6*               | 12.1±5.8                | 18.9±6.7†               |
| Epicardial stenosis resistance, mm Hg/cm per s | 0.26±0.06                  | 0.18±0.07             | 0.35±0.06               | 0.41±0.08               | 0.32±0.08               | 0.58±0.11*              | 0.33±0.08               | 0.46±0.09               |
| **Coronary hemodynamics**          |                             |                       |                          |                          |                          |                          |                          |
| Mean distal pressure, mm Hg        | 99.1±3.7                    | 114.5±3.7             | 87.7±10.9               | 102.6±11.8*             | 96.6±4.2                | 111.1±6.7*              | 100.6±3.8               | 121.8±7.8*               |
| Coronary blood flow velocity, cm/s | 18.1±1.7                    | 21.8±1.9*             | 17.3±2.1                | 16.8±2.2                | 19.9±2.7                | 32.1±4.3*               | 20.5±2.1                | 27.7±3.0               |
| Microvascular resistance, mm Hg/cm per s | 612±54                     | 576±48                | 550±44                  | 643±54*                 | 563±66                  | 397±43*                 | 550±42                  | 476±45               |
| Min. MVR, mm Hg/cm per s           | 449±43                      | 415±35                | 317±28                  | 394±28*                 | 373±47                  | 221±33*                 | 328±33                  | 288±40†               |
| **Coronary wave intensity analysis** |                             |                       |                          |                          |                          |                          |                          |
| Forward compression wave, J m⁻² s⁻³ 10⁵ | 0.52±0.09                  | 1.03±0.15*            | 0.59±0.09               | 0.56±0.12†              | 0.59±0.13               | 1.82±0.48*              | 0.56±0.10               | 1.56±0.42*               |
| Backward expansion wave, J m⁻² s⁻³ 10⁵ | 1.74±0.29                  | 2.98±0.36*            | 1.53±0.30               | 1.71±0.30†              | 1.91±0.21               | 6.47±1.34*              | 2.05±0.31               | 4.09±0.81†               |
| **Aortic hemodynamics**            |                             |                       |                          |                          |                          |                          |                          |
| Heart rate, bpm                    | 72.5±4.6                    | 76.3±3.6              | 74.1±3.4                | 75.1±2.9                | 79.4±3.8               | 121.9±5.5*              | 82.2±4.0                | 127.6±5.3*                |
| Mean aortic pressure, mm Hg        | 103.3±3.4                   | 117.9±3.7*            | 103.6±6.6               | 121.6±5.8*              | 107.0±3.0               | 131.7±4.2*              | 110.0±4.1               | 135.6±4.8*                |
| Systolic aortic pressure, mm Hg     | 143.5±5.0                   | 164.8±8.1*            | 139.4±10.1              | 166.6±8.1*              | 138.5±3.7               | 170.1±6.3*              | 141.6±4.6               | 181.5±7.0*                |
| Diastolic aortic pressure, mm Hg    | 72.8±3.4                    | 84.3±2.7*             | 78.0±4.7                | 87.4±5.0*               | 81.4±2.2                | 95.9±2.3*               | 82.7±2.7                | 97.5±3.1*                |
| Rate pressure product, mm Hg min⁻¹ 10⁶ | 10.4±0.8                    | 12.4±0.6*             | 10.4±0.9                | 12.5±0.8*               | 10.9±0.7               | 20.8±1.2*               | 11.5±0.4                | 23.1±1.1*               |
| Augmentation index, %               | 37.8±6.2                    | 45.8±5.2              | 38.7±6.0                | 49.9±10.4               | 51.1±10.4               | 27.2±6.1*               | 48.6±11.0               | 46.1±11.0               |
| Diastolic time fraction             | 0.59±0.02                   | 0.57±0.01             | 0.58±0.02               | 0.54±0.02               | 0.52±0.03               | 0.41±0.03*              | 0.53±0.02               | 0.35±0.02*†              |

Values are mean±SEM. Baseline indicates before application of cold air; Min. MVR, minimal MVR; obstructive stenoses, patients with obstructive coronary artery stenoses; Pa (mean), mean aortic pressure; Pa (mean), mean distal pressure; temp, temperature; U, coronary blood flow velocity; unobstructed coronaries, patients with unobstructed coronary arteries.

*P=0.05 vs respective baseline.
†P<0.05 vs peak exercise room temp.
‡P<0.05 vs peak normal coronary arteries.
Figure 5. Exercise with and without cold air inhalation. Change from baseline in patients with obstructive coronary stenoses, with the results subcategorized as per the following: (A) coronary hemodynamic results, (B) coronary wave intensity analysis results, and (C) aortic pulse wave analysis results.
Figure 6. Pathological effects of cold air during exercise in 12 patients with obstructive coronary stenoses. Ao indicates aorta; CBF, coronary blood flow velocity; LV, left ventricle; MVR, microvascular resistance; PA, pulmonary artery.

I. Exercise at Room Temperature – minor ischemia

1. More than two-fold increase in myocardial oxygen demand with exercise
2. Met by: 50% decrease in afterload, 30% decrease in microvascular resistance (MVR), and 60% increase in coronary blood flow velocity (CBF)
3. CBF increase principally driven by 200% increase in early diastolic CBF acceleration (shown by arrows)
4. This increase is enabled by effective myocardial relaxation at onset of diastole
5. Minor myocardial ischemia

II. Exercise during Cold Air Inhalation – greater ischemia

1. Similar increase in myocardial oxygen demand with exercise during cold air inhalation
2. But no reduction in afterload and only half the previous decrease in MVR and increase in CBF
3. Defective early diastolic CBF acceleration: less than half the previous increase (shown by smaller arrows)
4. Likely caused by failure of myocardial relaxation
5. Significantly more myocardial ischemia
The use of wave intensity analysis, unlike those of previous studies,24,25 has allowed us to examine the events within a single cardiac cycle that result in the mismatch between myocardial blood flow and demand. Our most striking finding was that in patients with obstructive coronary stenoses, cold air during exercise abrogated the adaptive 200% increase in the backward expansion wave that accompanied exercise in room temperature (Figure 6). This microcirculatory “suction” wave results from the release of external compressive forces on resistance vessels during ventricular relaxation,23 which in turn decreases MVR. This wave is the largest magnitude wave in the cardiac cycle and enables early diastolic coronary flow acceleration following aortic valve closure.23

Exercise in Room Temperature in Patients With Obstructive Stenoses

The mechanism of ventricular relaxation during exercise in room temperature deserves discussion. Despite increased myocardial oxygen demand and reduced diastolic time fraction,17 ischemia is minimized because of parallel adaptive reductions in afterload and MVR of 50% and 30%, respectively. The acute reduction in afterload enables a relatively efficient ejection time that is associated with enhanced ventricular relaxation,26 and in our study was evidenced by the 200% increase in the backward expansion wave. Ventricular relaxation during exercise may also be directly underpinned by efficient calcium reuptake via β-adrenoceptor-mediated protein kinase A activation.27 Enhanced ventricular relaxation reduces microvascular compression,28 which helps lower MVR. Despite vasoconstriction of the epicardial stenosed segment during exercise29 (evidenced by increased stenosis severity in our study), parallel intense vasodilatation of resistance vessels during exercise30 overcomes this, resulting in further adaptive reduction of MVR.

Cold Air During Exercise in Patients With Obstructive Stenoses

By contrast, ventricular relaxation is impaired with cold air during exercise (Figure 6). The adaptive reduction in afterload during exercise at room temperature is abolished: cold air caused a sufficiently potent sympathetic vasoconstrictive stimulus on the peripheral vasculature to offset the adaptive effects of dynamic exercise. This resulted in an additional pressure load on the ventricle, increasing relative ejection time and impairing ventricular relaxation.31 Ventricular relaxation may also be further impaired by relative autonomic conflict caused by parasympathetic activation of facial trigeminal receptors during cold air inhalation32: this would dampen β-adrenoceptor-mediated protein kinase A activation mentioned in the paragraph above. Furthermore, we observed a significantly exaggerated reduction in the diastolic time fraction during exercise with cold air. This is an important additional finding, because subendocardial blood flow has been shown to be dependent on diastolic time fraction, independently of heart rate.14 Parallel to the adverse effect on afterload, cold air during exercise also halved the adaptive reduction in MVR during exercise at room temperature. This is likely because of a combination of heightened α-1 adrenoceptor-mediated vasoconstriction exhausting the vasodilator reserve, systolic myocardial compression of resistance vessels because of increased ejection time, and impaired ventricular relaxation. These adverse effects may also be self-propagating, with increased myocardial ischemia further impeding ventricular relaxation via dysregulation of myocardial contractile machinery.

Unexpectedly, our measures of stenosis severity increased to a lesser extent during exercise in cold air versus exercise in room temperature (Table 2). This was seen in combination with a significantly higher peak minimal MVR during exercise in cold air versus exercise in room temperature. These data suggest that cold air during exercise results in relatively less epicardial resistance to coronary blood flow, but significantly more MVR to coronary blood flow. Therefore, these findings suggest that cold air–induced impaired vasodilatory capacity during exercise predominates at the arteriolar level, rather than the stenosed epicardial segment.

Cold Air Inhalation at Rest in Patients With and Without Obstructive Stenoses

MVR and coronary blood flow

Despite a similar increase in cardiac afterload to cold air inhalation, MVR decreased in patients with unobstructed coronary arteries but increased in those with obstructive coronary stenoses. While vasodilatation of small arteries and arterioles may prevail over α-1-adrenoceptor-mediated vasoconstriction in patients with unobstructed coronary arteries,33 patients with obstructive coronary stenoses are reliant on constant metabolic vasodilation of arterioles,9 evidenced by significantly lower minimal MVR at rest. These arterioles are therefore unable to compensate for vasoconstriction of small arteries in response to cold, resulting in increased MVR. In addition, these patients may have greater endothelial dysfunction and adverse structural remodeling of arterioles,34 resulting in a more potent vasoconstrictor response to cold air. This vasoconstrictor response predominantly affects the resistance vessels, not the stenosed epicardial artery, because we observed no significant increases in stenosis severity in either group (Table 2).

Wave intensity analysis

An interesting finding was the disparate response of the wave intensities to cold in patients with and without obstructive
coronary stenoses. Patients with unobstructed coronary arteries significantly increased the forward compression and backward expansion waves during cold air inhalation. This reflects rapid ventricular ejection and surge in left ventricular pressure, and is associated with efficient ventricular relaxation. However, patients with obstructive coronary stenoses likely have a reduced ventricular energetic reserve, which when combined with significant resistance vessel vasoconstriction predisposes the efficiency of ventricular ejection and relaxation to be compromised in the face of acutely increased afterload. In addition, downregulation of β-adrenoceptors following chronic sympathetic stimulation in patients with obstructive stenoses may also dampen the inotropic and chronotropic response to cold air. We found that the heart rate increase to cold air in patients with unobstructed coronary arteries was more than 3 times the magnitude than patients with obstructed coronary arteries. Therefore, at peak stress these patients were unable to increase either the forward compression or backward expansion wave.

Comparison With Other Studies

No previous studies have measured MVR, or applied wave intensity analysis or CMR perfusion techniques in response to cold air inhalation at rest or during exercise. Most studies for convenience used a “cold pressor test” at rest, whereby a limb is immersed into a bucket of iced water for 1 minute. However, this may not be physiologically relevant to exercising in cold air. In line with our findings, previous authors have found increases in MVR following a cold pressor test at rest using coronary sinus thermodilution in patients with coronary artery disease. This increase in MVR has been previously shown to result from vasoconstriction predominating at the arteriolar level, as the MVR increase was out of proportion to the small epicardial vasoconstrictor response demonstrated with quantitative coronary angiography. Also in response to cold pressor test in patients with coronary “luminal irregularities,” CBF measured with intracoronary Doppler increased by a similar magnitude to that observed in our patients with unobstructed coronary arteries. Literature investigating cold air during dynamic exercise is very sparse, with 1 invasive study demonstrating slightly smaller reductions in MVR during exercise with simultaneous cold pressor test, versus during exercise alone.

Mechanism of Increased Cardiac Mortality Caused by Cold Air During Exercise

Although the mechanism remains unclear, our study demonstrated significantly greater myocardial ischemia during exercise with cold air inhalation through a facemask versus exercise at room temperature. The burden of ischemia during exercise in the cold will likely be even greater with more skin exposure to cold, and/or a longer duration of cold exposure. Observations from long-distance running races suggest that ischemia burden alone may be sufficient to cause ventricular dysrhythmias and cardiac death. Moreover, cold stress may directly provoke arrhythmic cardiac death, through data demonstrating increased frequency of ventricular dysrhythmias during cold temperatures in patients with implanted cardioverter-defibrillators. Additionally, cold air inhalation may increase the likelihood of a plaque rupture event. Notably, in 9 patients presenting with acute myocardial infarction after shoveling snow, all had angiographic evidence of plaque rupture.

Unexpectedly, in our study we noted a smaller increase in trans-stenotic gradient during exercise in cold air versus room temperature. This would counterintuitively protect the stenosis against a shear stress and a plaque rupture event! However, this was associated with higher mean distal arterial pressure during exercise in cold air and potentially greater circumferential wall stress, arguably a more important predictor of plaque rupture. Shoveling snow and skiing involve isometric exercise in addition to dynamic exercise that will likely further increase myocardial oxygen demand and afterload.

Clinical Implications

These findings could change clinical practice. Doctors should inform patients with obstructive coronary stenoses about the following: (1) to avoid sudden vigorous exertion, as asymptomatic ischemia can manifest before angina occurs; (2) to avoid exertion in cold air where possible, or at least to wrap up warmly (ideally involving coverage of the cheeks, mouth, and forehead); (3) that their exercise tolerance may be less in cold air; (4) indoor warm-up exercise before exertion in cold air might attenuate these deleterious effects (by augmenting the reduction in MVR and increase in backward expansion wave intensity during the second period of exertion). In addition, angiotensin-converting enzyme inhibitors may attenuate coronary sympathetic vasoconstriction of resistance vessels, and glyceryl trinitrate may enhance diastolic ventricular relaxation, which may significantly reduce ischemia burden and possibly mortality with cold air.

Novel therapeutic approaches to preventing myocardial ischemia induced by cold air may attempt to preserve ventricular relaxation, which is determined by the clearance of the cytosolic calcium transient and the sensitivity of the myofilaments to calcium. Both these processes are thought to contribute to heart failure with preserved ejection and are the
subject of novel therapeutic intervention trials to influence them directly or indirectly.

Limitations
Our study has several important limitations. First, similar to other physiological studies investigating the effects of cold, the sample size was small, which limits extrapolation. Because this is a hypothesis-generating descriptive study, it is difficult to differentiate a association in a hemodynamic variable from another and we did not correct for multiple comparisons. Participants were predominantly male, and findings may not apply to females. Six ComboWire data sets were excluded from coronary hemodynamic and wave intensity analyses because of poor-quality traces. However, given the relatively homogeneous physiological responses in all variables, we believe our data are generally representative of our entire patient cohort.

Another weakness is the lack of a direct measure of ventricular relaxation or plasma catecholamines, although the latter have been measured before in response to cold. Further studies including intracoronary phentolamine and adenosine would be useful. We did not measure epicardial artery diameter changes during exercise because of safety concerns: Our protocol mandated removal of the guiding catheter away from the coronary ostium before starting exercise to reduce the risk of iatrogenic coronary artery dissection. Our control group of patients with unobstructed coronary arteries likely had endothelial dysfunction and impaired coronary vasomotion, because they were being investigated for chest pain with coronary angiography. However, this study was primarily designed to assess the effect of an obstructive coronary stenosis, and therefore our group classification remains relevant. Although we did not examine the effect of subjective cold air intolerance on the hemodynamic response to cold, cold tolerance was similar between groups. Finally, measurements of left ventricular pressure volume loops were not included in this exploratory study. Further studies might undertake simultaneous pressure volume loops and coronary wave intensity analysis.

Conclusions
In patients with obstructive coronary stenoses, cold air inhalation during exercise severely blunts the adaptive reductions in afterload and MVR that usually occur during exercise at room temperature. More detailed scrutiny by wave intensity analysis suggests that this is the result of a failure of coronary blood flow acceleration at the onset of diastole. In addition to the avoidance of exercising in cold air, this deficit may be amenable in the future to pharmacological manipulation.

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Disclosures
None.

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Supplemental Material
Data S1.

Supplemental Methods

Cardiac Magnetic Resonance (CMR) Myocardial Perfusion Sequence

Myocardial perfusion CMR was performed using dual bolus injection of
0.0075/0.075mmol/Kg of Gadolinium diethyl-enetriaminepentaacetic acid (Gadovist, Bayer Healthcare, Leverkusen, Germany) administered using a contrast saline
injector. Perfusion images were acquired in an end-expiratory breath-hold, with a high
spatial resolution saturation recovery gradient echo method (repetition time/echo time
3.0ms/1.0ms, flip angle 15°, 5-fold k-t broad linear speed up technique (BLAST)
acceleration and 11 training profiles, spatial resolution 1.2x1.2x10mm³, 90 dynamic
images). The mid-systolic slice, which was acquired in systole, was chosen for
analysis as this cardiac phase allows the myocardial perfusion layers to be more easily
separated.¹

Heart rate and blood pressure were measured continuously throughout each
protocol using a CMR compatible electrocardiogram and automated
sphygmomanometer (set to inflate automatically every minute) respectively.

CMR Perfusion Analysis

Subendocardial and subepicardial contours were delineated for every dynamic of the
perfusion series using dedicated software (EasyScil prototyping, Philips Medical
Systems, Best, Netherlands). The software generated signal-intensity curves for the
left ventricular blood pool and all myocardial segments, with myocardial signal-
intensity values corresponding to relative contrast concentrations from the first-pass
perfusion of Gadolinium. Perfusion estimates were obtained from signal-intensity
curves using a Fermi-function constrained deconvolution algorithm using Matlaboratory (MathWorks Inc., Natick, Massachusetts) as described by Jerosch-Herold et al. This generated absolute baseline and stress myocardial blood flow values (ml/g/min) for all 6 segments of the mid-ventricular slice.

Transmural perfusion gradients (TPG) were obtained using dedicated software (EasyScil prototyping, Philips Medical Systems, Best, Netherlands). Using the same contours delineated for deconvolution analysis, subendocardial, \textit{I}_{\text{endo}(a,t)}\), and subepicardial signal-intensity curves, \textit{I}_{\text{epi}(a,t)}\), were obtained. Ten myocardial layers and sixty radial segments were sampled and a signal-intensity curve generated at each point. TPG curves \textit{G}(a,t)\) were calculated based on the difference in subendocardial and subepicardial signal-intensity values over time, \textit{t} (each dynamic), at a particular myocardial location, \textit{a}, and normalized to transmural signal-intensity values, \textit{I}_{\text{transm}(a,t)}\),. Peak intensity of the TPG was expressed as a percentage of transmural flow redistribution. A high-pass threshold of TPG was set at 5% to reduce the effect of noise of the measured perfusion gradients. The radial extent of the TPG in angular degrees (°) was also calculated. A schematic representation of TPG analysis is shown in figure 2.

**Cardiac Catheterization Laboratory Data Analysis**

All pressure and flow signals were sampled at 200 Hz and stored on disk for off-line analysis. Ensemble averages of the selected cardiac cycles were performed for distal coronary artery pressure (\textit{P}_d) and coronary blood flow (estimated with average peak Doppler flow velocity (APV)). Savitzky–Golay filters were applied to preserve peaks in the pressure and flow data while smoothing. These filters fit a polynomial of a chosen order to a number of points around the centre point using least squares. They
have the advantage of smoothing data whilst preserving data peaks, which
dramatically improves the clinical applicability of coronary wave intensity analyses.

Net wave intensity (\(W_{\text{net}}\): W.m\(^{-2}\).s\(^{-2}\)) was calculated to be the product of time
derivatives (dt) of ensemble averaged and filtered coronary artery pressure (Pd) and
flow (U) signals:\(^6\)

\[
\frac{\Delta I_{\text{net}}}{\Delta t} = \frac{\Delta Pd}{\Delta t} \cdot \frac{\Delta U}{\Delta t}
\]

\(W_{\text{net}}\) can be separated into forward contributing and backward components
arriving at the measurement site:\(^7\)

\[
\frac{\Delta I_{\text{net}}}{\Delta t} = \Delta I_+ + \Delta I_-
\]

Where forward travelling waves (WI\(_+\)) are defined by a positive WI, which
occurs when dP\(_d\) and dU change in the same direction. In contrast backward travelling
waves (WI\(_-\)) occur when dP\(_d\) and dU change in opposite directions, leading to a
negative WI.

Waves are also defined as accelerating when dU increases, and decelerating
when dU decreases. In addition waves can be defined as compression waves when
associated with an increase in dP\(_d\), or defined as expansion waves when associated
with a decrease in dP\(_d\). Four dominant waves were calculated: the accelerating
forward compression wave (FCW); the decelerating forward expansion wave (FEW);
the decelerating backward compression wave (BCW); and the accelerating backward
expansion wave (BEW).\(^8\) In the coronary circulation forward travelling waves are
generated by increases (FCW) and decreases (FEW) in aortic pressure at the inlet and
backward travelling waves are generated by changes in the microcirculation due to
cardiac contraction (BCW) and relaxation (BEW).\(^9-11\) Of these four waves, the
acceleratory waves produced the greatest magnitude. Therefore only the two
acceleratory waves, FCW and BEW, are reported in the manuscript.
In order to separate WI into the forward (WI+) and backward (WI-) components these respective formulae were applied to collected data:

\[ WI_+ = \frac{1}{4\rho c^2} \left( \frac{\partial P}{\partial t} + \rho \frac{\partial U}{\partial t} \right)^2 \]

\[ WI_- = -\frac{1}{4\rho c^2} \left( \frac{\partial P}{\partial t} - \rho \frac{\partial U}{\partial t} \right)^2 \]

Where \( \rho \) = density of blood, \( c \) = wavespeed, and WI was defined as the product of the first time derivatives of coronary artery pressure and flow velocity so that the analysis is independent of the sampling interval used, as previously described.\(^{12}\) The density of blood was assumed to be constant at 1050kgm\(^{-3}\).\(^7\) Wavespeed was calculated using the single-point technique as previously described.\(^7\)

Rate pressure product (RPP), a measure of myocardial oxygen demand,\(^{13}\) was calculated as systolic blood pressure multiplied by heart rate. Diastolic time fraction was defined as the fraction of the duration of diastole with respect to duration of the cardiac cycle. Systolic duration was defined from the upslope of the arterial pressure trace to the dichrotic notch, and diastolic duration defined from the dichrotic notch to the upslope of the arterial pressure trace. Augmentation index, a measure of central systolic blood pressure augmentation thought to principally arise from pressure-wave reflection, was calculated as the difference between the first and second aortic systolic pressure peaks expressed as a percentage of the pulse pressure (in turn calculated as systolic blood pressure minus diastolic blood pressure).
**Figure S1. Cardiac Catheterization Laboratory Protocol Set Up.**

**A.** For cold air inhalation we adapted a clinical cryotherapy device with a closed cooling circuit: Cryo6 Cold Air Device (Zimmer Medizin Systeme, Ulm, Germany), at a fixed temperature of 5°F and a flow rate of 50L/min. A standard non-rebreathe facemask was adapted and fitted onto the end of the nozzle of the cryotherapy tubing. This therefore provided a means of supplying cold air for cold air inhalation.

**B.** Facemask application. The cold air chilled the cheeks, nose, mouth and chin, and mouth breathing was encouraged where possible (to avoid excess nasal warming of cold air).

**C.** An intracoronary 0.014-in guidewire (ComboWire, Volcano® Corporation, San Diego, USA) with Doppler velocity transducer on the wire tip and pressure transducer 1.5cm proximally to this on the wire shaft. This enabled simultaneous and continues measurement of coronary artery flow velocity and pressure. The ComboWire was inserted into the distal coronary artery (beyond the coronary artery stenosis in patients with coronary artery disease). When feasible the wire was inserted into the vessel retrogradely on a loop, which improved stability of the Doppler signal.

**D.** ComboMap console (Volcano® Corporation, San Diego, USA) demonstrating simultaneous acquisition of aortic pressure waveforms (red), distal coronary artery pressure waveforms (yellow) and Doppler velocity (blue with underlying grey scale) waveforms.
**Figure S2. Transmural Perfusion Gradients with CMR Perfusion Scans.**

The grey scale images on the top half of this figure demonstrate typical high-resolution CMR perfusion images of three short-axis slices of the left ventricle. In this example there is visible subendocardial ischemia in the mid-ventricular and basal slices (shown as a darker rim of grey scale on the inside of the myocardium, highlighted with *). In order to quantify transmural perfusion gradients (TPG), subendocardial, subepicardial and left ventricular blood pool contours are drawn onto these images using EasyScil prototyping software. The TPG algorithm calculates the intensity of the gradient in each angular and temporal position by the spatial averaging of the signal intensity (of gadolinium contrast in the myocardium) of the inner and outer third of the left ventricular wall, normalised by the average transmural signal intensity to account for signal inhomogeneity. This creates a gradientogram plot that is shown on the lower half of this figure. The amplitude of the TPG is represented by the intensity of grey scale colour, so that a darker area represents a region of poorer subendocardial perfusion. A threshold TPG % can be set whereby an area above this set threshold represents a subendocardial perfusion deficit, as highlighted in green and highlighted with * (corresponding to the visual CMR perfusion images above). Temporal persistence in seconds (the length of time that the TPG is apparent for) is measured along the x-axis, and circumferential extent in angular degrees (the extent of myocardium involved) is measured along the y-axis.

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Figure S3. Transmural Perfusion Gradients during Cold Air Inhalation in Patients with Significant Coronary Artery Disease.

We observed significant increases in peak TPG intensity and radial TPG extent (both p<0.05). Peak TPG intensity reflects the subepicardial-subendocardial redistribution of myocardial blood flow, and radial extent TPG reflects the amount of myocardium affected. Increases in both parameters suggest an insufficiency of subendocardial perfusion. A peak TPG intensity value of greater than 20% (during adenosine hyperemia) has been shown to strongly correlate with a significant fractional flow reserve (<0.8).\textsuperscript{14}
Supplemental Video Legend:

Video S1. Cardiac Catheterization Laboratory Protocol Set Up.
Supplemental References:

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